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### (12) United States Patent

#### Bandara et al.

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# (54) DEVELOPMENT OF A LIVE, ATTENUATED, RECOMBINANT VACCINE FOR BRUCELLOSIS

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(US)

# (73) Assignee: Virginia Tech Intellectual Properties, Inc., Blacksburg, VA (US)

# (\*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35

U.S.C. 154(b) by 111 days.

This patent is subject to a terminal dis-

claimer.

(21) Appl. No.: 11/246,957

(22) Filed: Oct. 11, 2005

### (65) Prior Publication Data

US 2006/0093621 A1 May 4, 2006

#### Related U.S. Application Data

- (63) Continuation-in-part of application No. 11/050,429, filed on Feb. 4, 2005.
- (60) Provisional application No. 60/541,954, filed on Feb. 6, 2004.
- (51) **Int. Cl.**A61K 39/10 (2006.01)
  C12N 15/74 (2006.01)
- (52) **U.S. Cl.** ...... **424/252.1**; 435/471; 435/477
- (58) **Field of Classification Search** ....... None See application file for complete search history.

#### (56) References Cited

#### U.S. PATENT DOCUMENTS

5,718,903	A *	2/1998	Adams et al 424/235.1
5,851,519	A *	12/1998	Dougan et al 424/93.2
5,939,075	A *	8/1999	Houng et al 424/252.1
6,149,920	A *	11/2000	Boyle et al 424/252.1
6,296,855	B1 *	10/2001	Hemmen et al 424/252.1
6,444,445	B2 *	9/2002	Nikolich et al 435/69.3
6,582,699	B2 *	6/2003	Cherwonogrodzky 424/184.1

#### (Continued)

#### FOREIGN PATENT DOCUMENTS

ES 2140336 \* 2/2000

#### (Continued)

#### OTHER PUBLICATIONS

DelVecchio et al., 2002, "The genome sequence of the facultative intracellular pathogen *Brucella melitensis*", Proceedings of the National Academy of Sciences, USA, vol. 99, No. 1, pp. 443-448.\*

#### (Continued)

Primary Examiner—Nashaat T. Nashed Assistant Examiner—William W. Moore (74) Attorney, Agent, or Firm—Whitham Curtis Christofferson & Cook, PC

### (57) ABSTRACT

A recombinant, attenuated strain of *Brucella suis* or *Brucella melitensis* with a deficiency in carboxyl-terminal protease activity or tail-specific protease activity can be used as a vaccine for the prevention or treatment of Brucellosis. Prior exposure to the *Brucella* species is identified by detecting a genetic sequence for carboxyl-terminal (i.e. tail-specific) protease activity in a biological sample.

#### 7 Claims, 16 Drawing Sheets

36 TRODDEKLVENAINGHLSSLOPKSSFMNAKDANDMRTDTKGBFGGLGISVTHE	
53 TIPDDOKLIENAINGKLLSLDPKSSYNDAEFAKDMRDSTXGEFGGLGIEVTME	
60 IKPDNAKLIEGAITGKVTSLDPKSRYMNDKAWTEMQETTSGEFGGLGIEVTME	
59 TPPDDKKLIESAINGHLTSLDPKSSYLNPEAAQDMRVQTXGEDGGLGIEVTH	DNDLVKVI Brucell
57 TPPDDKSLVENAINGMLSSLDPKSSYMNAEGAQDMRVQTKGEFGGLGIEVTME	ENDLVKVI Mesorhi
61 EKPDDSKLVESAISGKLAGLDPKSSYMDAKSFRDMQVQTRGEFGGLGIEVTM	
56 TPPQDDKLIENAINGKLTSLDPKSSYMNSTDAEDMRTQTRGEFGGLGIEVTMS	SECLVKVT Simorhi
11/ 02/12/04/14 14/14/14/14/14/14/14/14/14/14/14/14/14/1	
116 SPMDDTPASRAGILAGDYISBIDGTPVRGLKLEQAVEKMRGAVKTPIKLTVIF 113 SPIDDTPAAKAGVLAGDFISKIDGKQISGQTLMBAVDQMRGPAGTPITLTIM	
120 SPIDDTPASKAGIKSGDLISKIDGDAVQGMTLEQAVMKMRGPVDTKTKLTIV	
119 APIDDTPASKAGVLSGDLITKIDGQEVRGLSLPDAVDKMRGEVGAPIELTILI	
110 TPIDDTPAAKAGVLAGDYIAKIDGKEVRGLTLNDAVDKMRGLVNTPIKLTIL	
121 SPIDDTPASKAGILANDIITNIDDEAVQGLTLNQAVEKMRGPVNTKIALKIV	REGUDNET Rhodops
136 SPIDDTPAARAGVLAGDFISKIDGQDVRGLKLESAVDKMRGAVGTPIKLTILE	RKGADKP1 Sinorhi
176 EPTVVRDVJAVRAVKSRVEGDNVGYLRVISPTEKTYDDKEKAIKKIKADVPAL	NELECTVI Agrabia
173 DIKIVRDIIKVKAVKYRVEGD-IGYLRLIQFTEKTFSDLQAAIKDIQSKIPTI	
180 DIAJTRDIIHVRPVRPNVBNGDIGYIRVTSPNEQT7DGLKKAIAAISREIPQS	EKLAGVVK Bradyrh
179 ELKJNRAIIKVKAVRSRVEND-VGYLRIISPPEQTSEDLKKAIKDIQEKVPA	
177 ELTVVRDIIKVKAVKPRVEND-IGYMRITSPTEKTYDDLENAIDPIKKQVPDI	
181 EVTLVRDNIRVRSVRARVEDSDIGYIRITTPNEQTTEGLRKEIANLTNQIGA	
176 ELTIVEDVIAVEAVEVEVEGD-VGYLEVISPTEKTPSDLKEGIEKIQAEVPA	
JUL	
235 DLRLNPGGLLDQAINVSDAPLERGEVVSTRGRNPDETRRPNATAGDLTDG	
232 DLRLNPGGLLDQAISVTDAPLNKGEIVSTRGRKQNDVNRPDAKLGDLTDE	
210 DLRNNPGGLLDQAVSVSSAPLQRGEVVSTRGRNPBETQRPTANGGDLTKG	
238 DLRLNPGGLLDQAVAVSDAPLDKGEIVSTRGRDPQDVTRPDVRNGDLTNG	
236 DLRLNPGGLLDQAVSVSDAPLKRGEIVSTRGRDPKDVRRPDALPSQTDDING	
241 DLRNNPGGLLEEAVTVSDAPLDRGEIVSTRGRNAEETQRRDARAGDLTRG	
235 DLRLNPGGLLDQAINVSDAPLETGBVVSTRGRNPDETRRPNAT?GDLAGG	(PVVVLVN Sinorhi
294 GGSASASEIVAGALQDLRRATVVGTRSPGKGSVQTIIPLG-BAGALRITTAL	VVTDSCKS Larabia
290 GGSASASEIVAGALQDRRRATIIGTQSPGKGSVQTIIPLG-ENGALRITTALY	YYTPSGTS Bartone
298 GGSASASEIVAGALRORKRATIIGTRSPGKGSVQTIIPLGAGNGALALTTAR	YYTPSGES Bartone
296 GGSASASBIVAGALQDRRRATVLGTOSPGKGSVOTIIPLG-ENGSLRLTTAL)	
298 GGSASASBIVAGALODRRRVTVVGTOSPGKGSVOTIIPLG-ZNGALRLTTAL)	
299 GGSASASBIVAGALQDRXRATVVGTRSPGKGSVQTIIPLGSGKGALRLTTAR	
293 GGSASASBIVAGALQDLKRATVLGTRSPGKGSVQTIIPLG-DAGALRLTTAL	
TO GOSSON DE LEGENA DE MANTEN DE LA DECENSA DE LE LES DE CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DE LA CONTRA DE LA CONTRA DE LA CONTRA DE	iiirauna Sinorni

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### U.S. PATENT DOCUMENTS

6,811,787 B1*	11/2004	Boyle et al 424/252.1
2003/0044431 A1*	3/2003	Schurig et al 424/252.1
2005/0142151 A1*	6/2005	Nikolich et al 424/252.1
2005/0249755 A1*	11/2005	Nikolich et al 424/252.1
2006/0153868 A1*	7/2006	Ugalde et al 424/200.1

#### FOREIGN PATENT DOCUMENTS

WO WO 88/07374 \* 10/1988

WO WO 98/08951 \* 3/1998

### OTHER PUBLICATIONS

Paulsen et al., 2002. "The *Brucella suis* genome reveals fundamental similarities between animal and plant pathogens and symbionts", Proceedings of the National Academy of Sciences, USA, vol. 99, No. 20, pp. 13148-13153.\*

\* cited by examiner

teagtt cageacgeee tttttegggt eeggegggaa ageggeattg ggeaettege eeegcagaag cttcagtgcc tcattgagct ggagatcatc cttcggatca ggcggaacat aagccgacga accggageeg etegeatett eggeattgee ettgatatgg eeettgagtt eggattegee gegaacaaeg teetegeett taagtteegg eggeagegge tgateeaeet tgatgteegg cgtaatgccc ttgccctgga tcgacttgcc agacggcgtg taataaagcg ccgtcgtcag acgeagegaa cegittiege caagegggat gattgietge acagagecet tgeegaagga ctgcgtaccg agcaccgtgg cgcggcgatg atcctgaagc gcaccggcaa cgatttccga ggcactggcc gaaccgccat tgatcagaac gatcageggc ttgccattcg tcaggtcacc cttacgggca tcgaaacggg tcacatcctg cggatcgcgg ccacgggtgg aaacgatctc gecettgteg aggaaggeat eggaaaegge aacegeetga teeagaagge egeeeggatt gagacgaagg tegageacat agecettgag ettgteageg ggaacetttt eetgaatgte cttgategee ttettgaggt etteagaggt etgtteggta aacgagatga taegeagata gecaacatea tteteaaege gegagegaae egeetteaee ttgataatgg egegattgat ettgagegtg ateggettgt eggegeeett gegeaggate gteagttega ttggegeace aacctegeeg egeatettgt ccacagegte ggteagegaa aggeegegea etteetgtee gtcgatcttg gtgatcaggt egecegacag cacaceggee ttegaggegg gegtategte gateggegeg atgacettea aaactcaccc ttggtctgca egagateatt gteeatggtg aettegatge caaggeegee cacgcatgtc ctgcgcggct tccgggttga gataggacga atggggatca agagaagtca geatgeeatt gatggegete tegateaget tettgtegte gggcggcgtc acatattgcg egegtacaeg etegaaaata tegeegaaaa gageeagete cttatagaca tegetgtett tteetgeege aaaageggtg gaagetggtg egeeetggae cateaceat

# Figure 1A

MVMVQGAPASTAFAAGKDSDVYKELALFGDIFERVRAQYVTPPDDK KLIESAINGMLTSLDPHSSYLNPEAAQDMRVQTKGEFGGLGIEVTMD NDLVKVIAPIDDTPASKAGVLSGDLITKIDGQEVRGLSLTDAVDKMR GEVGAPIELTILRKGADKPITLKINRAIIKVKAVRSRVENDVGYLRIISF TEQTSEDLKKAIKDIQEKVPADKLKGYVLDLRLNPGGLLDQAVAVSD AFLDKGEIVSTRGRDPQDVTRFDARKGDLTNGKPLIVLINGGSASASE IVAGALQDHRRATVLGTQSFGKGSVQTIIPLGENGSLRLTTALYYTPS GKSIQGKGITPDIKVDQPLPPELKGEDVVRGESELKGHIKGNAEDAS GSGSSAYVPPDP KDDLQLNEALKLLRGEVPNAAFPPDPKKGVLN

## Figure 1B

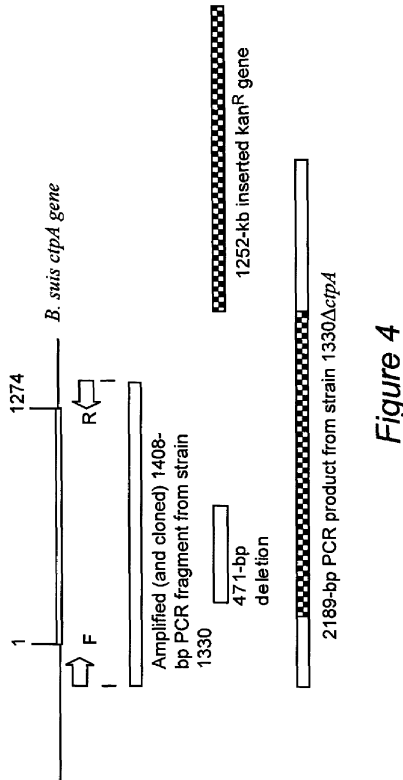
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# Figure 2

teagtt cageaegeee tttttegggt eeggegggaa ageggeattg ggeaettege eeegeagaag etteagtgee teattgaget ggagateate etteggatea ggeggaacat aageegaega accggagccg ctcgcatctt cggcattgcc cttgatatgg cccttgagtt cggattcgcc gegaacaaeg teetegeett taagtteegg eggeagegge tgateeaeet tgatgteegg cgtaatgccc ttgccctgga tcgacttgcc agacggcgtg taataaagcg ccgtcgtcag acgcagcgaa cegttttege caagcgggat gattgtetge acagagccet tgccgaagga ctgcgtaccg ageaccgtgg cgcggcgatg atcctgaagc gcaccggcaa cgatttccga gaaccgccat tg=====deletion\*====== ggcactggcc atcaggt cgcccgacag cacaccggcc ttcgaggcgg gcgtatcgtc gatcggcgcg atgaccttca aaactcaccc ttggtctgca cgagatcatt gtccatggtg acttcgatgc caaggccgcc cacgcatgtc ctgcgcggct tccgggttga gataggacga atggggatca agagaagtca geatgeeatt gatggegete tegateaget tettgtegte gggcggcgtc acatattgcg egegtacaeg etegaaaata tegeegaaaa gageeagete cttatagaca tcgctgtctt ttcctgccgc aaaagcggtg gaagctggtg cgccctggac catcaccat

\* A gene encoding resistance to antibiotic Kanamycin was incorporated to this deleted region.

# Figure 3



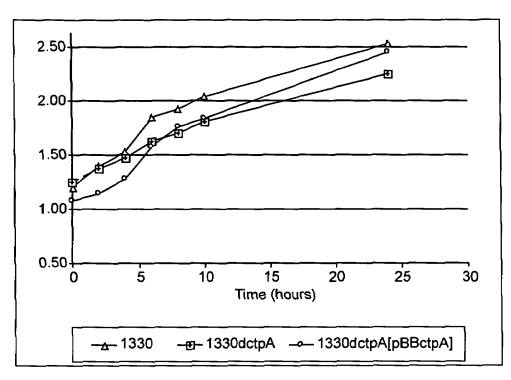


Figure 5A

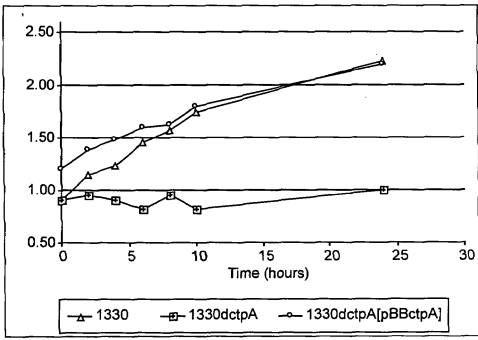


Figure 5B

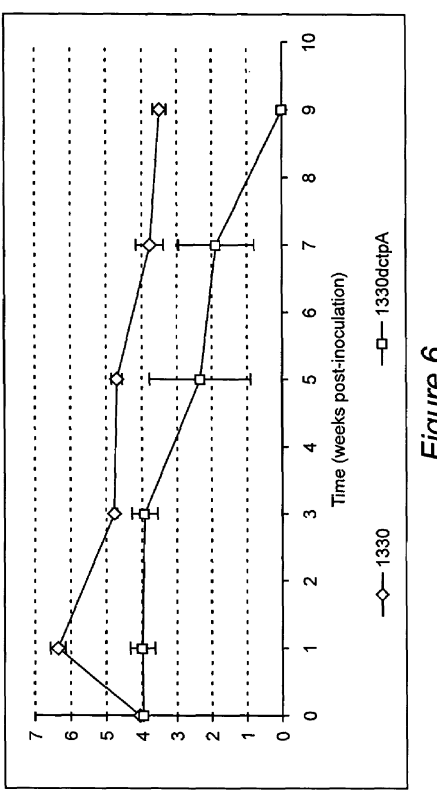


Figure 6

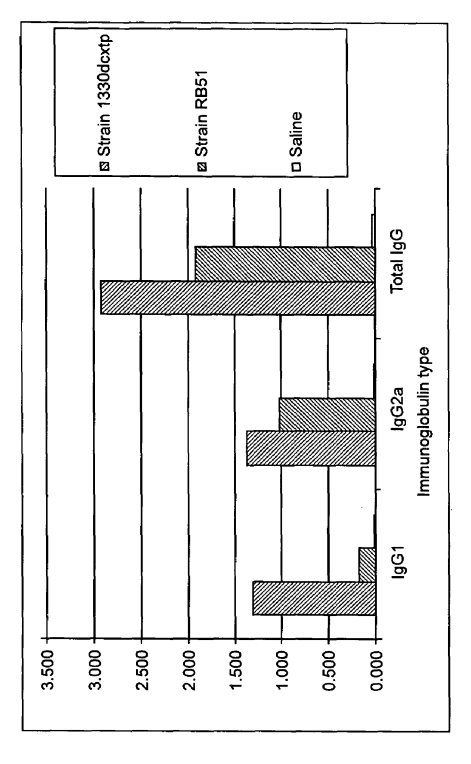


Figure 7

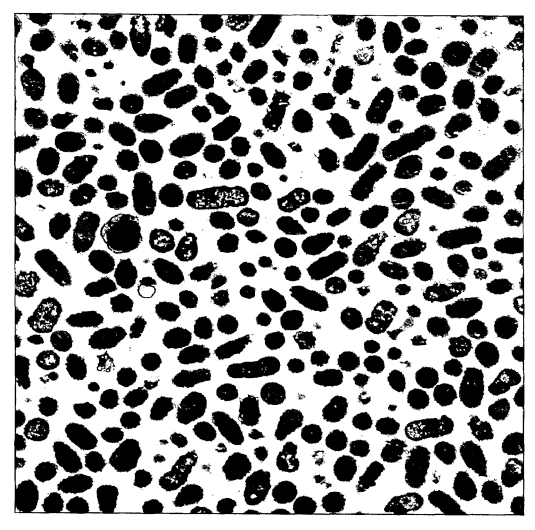


Figure 8

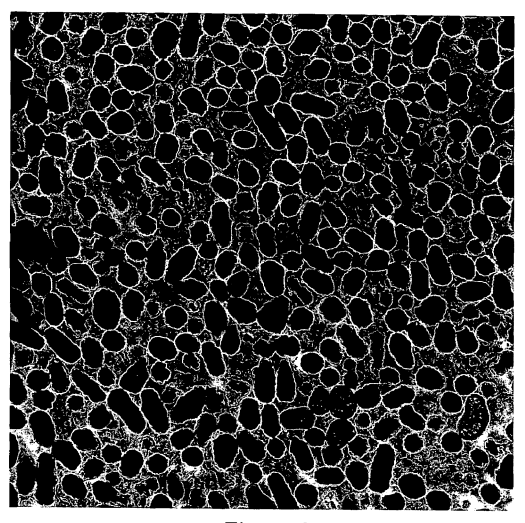


Figure 9

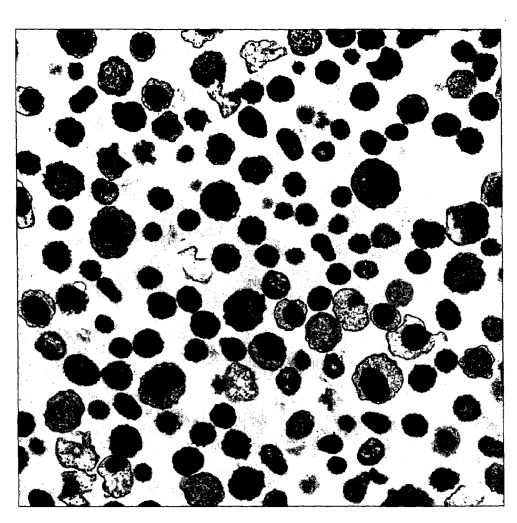


Figure 10

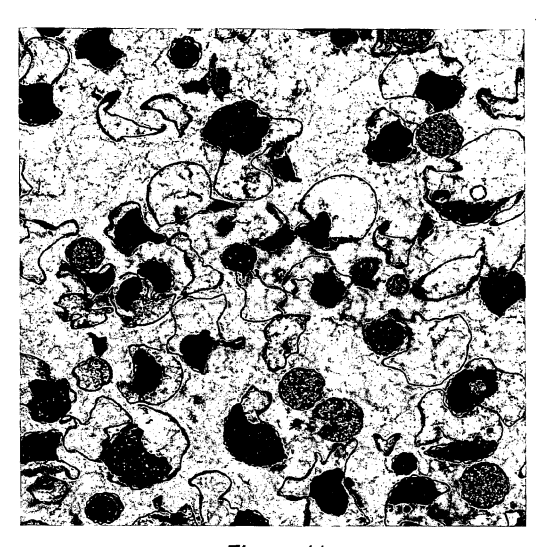


Figure 11

atgatacgtaaactgtcgctgctgttcgccggggcccttctgggggcatccgccatggtgatggtccagggggcaccagc ttccaccgcttttgcggcaggaaaagacagcgatgtctataaggagctggctcttttcggcgatattttcgagcgtgtac gegegeaatatgtgaegeegeegaegaeaagaagetgategagagegeeateaatggeatgetgaettetettgateee cattogtcctatctcaacceggaagcegegeaggacatgegtgtgcagaccaagggtgagtttggeggecttggcatega agteaccatggacaacgatetegtgaaggteategegeegategacgatacgeeegeetegaaggeeggtgtgetgtegg gegacetgateaceaagategaeggaeaggaagtgegeggeetttegetgaeegaegetgtggaeaagatgegeggegag gttggtgcgccaatcgaactgacgatcctgcgcaagggcgccgacaagccgatcacgctcaagatcaatcgcgccattat caaggtgaaggcggttcgctcgcgcgttgagaatgatgttggctatctgcgtatcatctcgtttaccgaacagacctctgaagacctcaagaaggcgatcaaggacattcaggaaaaggttcccgctgacaagctcaagggctatgtgctcgaccttcgt cteaateegggeggeettetggateaggeggttgeegttteegatgeetteetegaeaagggegagategttteeaeeeg tggccgcgatccgcaggatgtgacccgtttcgatgtccgtaagggtgacctgacgaatggcaagccgctgatcgttctga teaatggeggtteggeeagtgeeteggaaategttgeeggtgegetteaggateategeegegeeaeggtgeteggtaeg cagtcetteggeaagggetetgtgeagacaateateeegettggegaaaaeggttegetgegtetgaegaeggegettta ttacacgccgtctggcaagtcgatccagggcaagggcattacgccggacatcaaggtggatcagccgctgccgccggaac ttaaaggegaggacgttgttegeggegaateegaacteaagggeeatateaagggeaatgeegaagatgegageggetee ggttcgtcggcttatgttccgcctgatccgaaggatgatctccagctcaatgaggcactgaagcttctgcggggcgaagt ggccaatgccgctttcccgccggacccgaaaaagggcgtgctgaactga

# Figure 12A

MIRKLSLLFAGALLGASAMVMVQGAPASTAFAAGKDSDVYKELALFGDIFERVR AQYVTPPDDKKLIESAINGMLTSLDPHSSYLNPEAAQDMRVQTKGEFGGLGIEVT MDNDLVKVIAPIDDTPASKAGVLSGDLITKIDGQEVRGLSLTDAVDKMRGEVGA PIELTILRKGADKPITLKINRAIIKVKAVRSRVENDVGYLRIISFTEQTSEDLKKAIK DIQEKVPADKLKGYVLDLRLNPGGLLDQAVAVSDAFLDKGEIVSTRGRDPQDVT RFDVRKGDLTNGKPLIVLINGGSASASEIVAGALQDHRRATVLGTQSFGKGSVQT IIPLGENGSLRLTTALYYTPSGKSIQGKGITPDIKVDQPLPPELKGEDVVRGESELK GHIKGNAEDASGSGSSAYVPPDPKDDLQLNEALKLLRGEVANAAFPPDPKKGVL N

Figure 12B

```
36 TRDDDEKLVENAINGMLSSLDPKSSFMNAKDANDMRTDTKGEFGGLGISVTMENELVKVI Agrobic
53 TIPDDOKLIENAINGKLLSLDPKSSYNDAEFAKDMRDSTXGEFGGLGIEVTMENNLIKVV Bartone
60 IKPDNAKLIEGAITGKVTSLDPKSRYMNDKAWTEMQETTSGEFGGLGIEVTMEEGLVKVV Bradyrh
59 TPPDDKKLIESAINGMLTSLDPKSSYLNPEAAQDMRVQTXGEDGGLGIEVTMDNDLVKVI Brucell
57 TPPDDKSLVENAINGMLSSLDPKSSYMNAEGAQDMRVQTKGEFGGLGIEVTMENDLVKVI Mesorhi
61 EKPDDSKLVESAISGMLAGLDPKSSYMDAKSFRDMQVQTRGEFGGLGIEVTMEDGLIKVV Rhodops
56 TPPQDDKLIENAINGKLTSLDPKSSYMNSTDAEDMRTQTRGEFGGLGIEVTMSEDLVKVT Sinorhi
116 SPMDDTPASRAGILAGDYISEIDGTPVRGLKLEQAVEKMRGAVKTPIKLTVIRKGADKPL Agrobic
113 SPIDDTPAAKAGVLAGDFISKIDGKQISGQTLNEAVDQMRGPAGTPITLTINRFGVDKPL Bartone
120 SPIDDTPASKAGIKSGDLISKIDGDAVQGMTLEQAVMKMRGPVDTKTKLTIVRKGADAPL Bradvrh
119 APIDDTPASKAGVLSGDLITKIDGQEVRGLSLPDAVDKMRGEVGAPIELTILRKGADKPI Brucell
110 TPIDDTPAAKAGVLAGDYIAKIDGKEVRGLTLNDAVDKMRGLVNTPIKLTILRQGADKPL Mesorhi
121 SPIDDTPASKAGILANDIITNLDDEAVQGLTLNQAVEKMRGPVNTKIALKIVRKGQDNPI Rhodops
136 SPIDDTPAARAGVLAGDFISKIDGQDVRGLKLESAVDKMRGAVGTPIKLTILRKGADKP1 Sinorhi
176 EPTVVRDV JAVRAVKSRVEGDN VGYLRVISPTEKTYDDKEKAIKKIKADV PADKLKGVVL Agrobic
173 DIKIVRDIIKVKAVKYRVEGD-IGYLRLIQFTEKTFSDLQAAIKDIOSKIPTDKLKGVVL Bartone
180 DIAJTRDIIHVRPVRPNVENGDIGYIRVTSPNEQTTDGLKKAIAAISREIPQEKLAGVVK Bradyrh
179 ELKJNRAIIKVKAVRSRVEND-VGYLRIISPPEQTSEDLKKAIKDIQEKVPADKLKGVVL Brucell
177 ELTVVRDIIKVKAVKPRVEND-IGYMRITSPTEKTYDDLENAIDPIKKQVPDDKLKGVV; Mesorhi
181 EVTLVRDNIRVRSVRARVEDSDIGYIRITTPNEQTTEGLRKEIANLTNQIGADKLKGPIL Rhodops
176 ELTIVRDVIAVRAVKVRVEGD-VGYLRVISPTEKTPSDLKKGIEKIQAEVPADKLKGYVL Sinorhi
235 DLRLNPGGLLDQAINVSDAPLERGEVVSTRGRNPDETRRPNATA--GDLTDGKPVIVLVN Agrobic
232 DLRLNPGGLLDQAISVTDAPLNKGEIVSTRGRKQNDVNRPDA--KLGDLTDEKPIIVLIN Bartone
210 DLRNNPGGLLDQAVSVSSAPLQRGEVVSTRGRNPEETQRPTANG--GDLTKGKPLVVLVN Bradyrh
238 DLRLNPGGLLDQAVAVSDAPLDKGEIVSTRGRDPQDVTRPDVR--NGDLTNGKPLIVLIN Brucell
236 DLRLNPGGLLDQAVSVSDAPLKRGEIVSTRGRDPKDVRRPDALPSQTDDINGKPMIVLVN Mesorhi
241 DLRNNPGGLLEEAVTVSDAPLDRGEIVSTRGRNAEETQRRDARA--GDLTRGKPVIVLIN Rhodops
235 DLRLNPGGLLDQAINVSDAPLETGEVVSTRGRNPDETRRPNATP--GDLAGGKPVVVLVN Sinorhi

I
290 GGSASASEIVAGALQDRRRATIIGTQSPGKGSVQTIIPLG-ENGALRLTTALYYTPSGTS Bartone
298 GGSASASEIVAGALRDRKRATIIGTRSPGKGSVQTIIPLGAGNGALALTTARYYTPSGRS Bradyrh
296 GGSASASEIVAGALQDRRRATVLGTQSPGKGSVQTIIPLG-ENGSLRLTTALYYTPSGKS Brucell
298 GGSASASEIVAGALQDRRRVTVVGTQSPGKGSVQTIIPLG-ENGALRLTTALYYTPSGKS Mesorhi
299 GGSASASEIVAGALQDRKRATVVGTRSPGKGSVQTIIPLGSGNGALRLTTARYYTPSGKS Rhodops
293 GGSASASEIVAGALQDLKRATVLGTRSPGKGSVQTIIPLG-DAGALRLTTALYYTPSGKS Sinorhi
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Figure 13

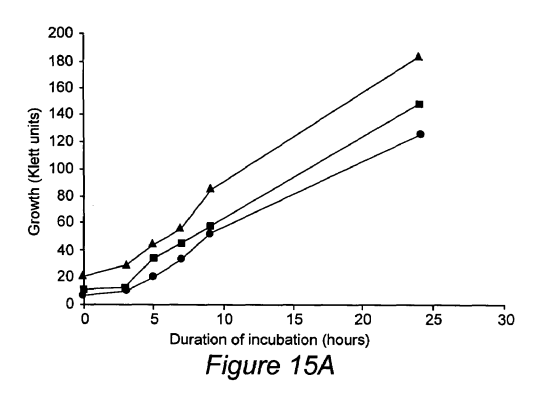
# Figure 14A

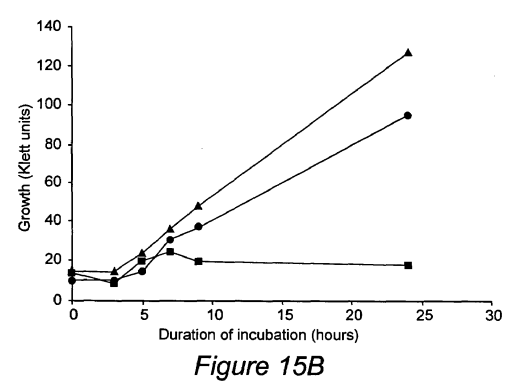
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### Figure 14B

gatcacca agatcgacgg acaggaagtg cgcggccttt cgctgaccga cgctgtggac aagatgcgcg gcgaggttgg tgcgccaatc gaactgacga tcctgcgcaa gggcgccgac aagccgatca cgctcaagat caatcgcgcc attatcaagg tgaaggcggt tcgctcgcg gttgagaatg atgttggcta tctgcgtatc atctcgttta ccgaacagac ctctgaagac ctcaagaagg cgatcaagga cattcaggaa aaggttcccg ctgacaagct caagggctat gtgctcgacc ttcgtctcaa tccgggcggc cttctggatc aggcggttgc cgtttccgat gccttcctcg acaagggcga gatcgtttcc acccgtggcc gcgatccgca ggatgtgacc cgtttcgatg tccgtaaggg tgacctgacg aatggcaagc cgctgatcgt tct

# Figure 14C





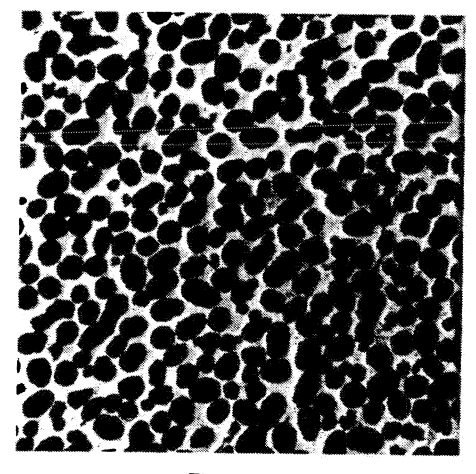


Figure 16A

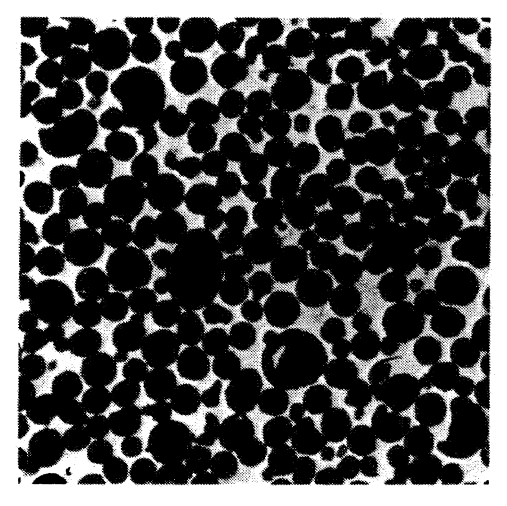


Figure 16B

#### DEVELOPMENT OF A LIVE, ATTENUATED, RECOMBINANT VACCINE FOR BRUCELLOSIS

This application claims benefit of U.S. provisional patent 5 application U.S. 60/541,954, filed Feb. 6, 2004, and is a continuation-in-part of U.S. patent application Ser. No. 11/050,429, filed Feb. 4, 2005, the complete contents of each of which are hereby incorporated by reference.

This invention was made using funds from grants from  $_{10}$  the United States Department of Agriculture having grant number 1-37181. The United States government may have certain rights in this invention.

#### BACKGROUND OF THE INVENTION

#### 1. Field of the Invention

The invention generally relates to a vaccine for Brucellosis. In particular, the invention provides an antigenic composition comprising a recombinant, attenuated strain of *Brucella suis* with a deficiency in carboxyl-terminal protease <sup>20</sup> (CtpA) activity.

#### 2. Background of the Invention

Animal brucellosis is a disease affecting many domestic and wild life species. In male animals, this disease causes orchitis (inflammation of the testicles) and may eventually 25 lead to sterility. In female animals, brucellosis causes abortion during the last trimester, retained afterbirth (retaining placenta in the uterus) and weakness in calves at birth. Brucellosis results from infection with bacteria belonging to the genus Brucella. On the basis of observed differences in 30 host preference, which have been associated with certain phenotypic characteristics, this genus has been classified for convenience into six nomen species. These are associated with different principal hosts: B. abortus (cattle), B. canis (dogs), B. melitensis (sheep, goats), B. neotomae (desert wood rat), B. ovis (sheep) and B. suis (swine, reindeer). However, Brucella species typically can infect a wide variety of hosts, including humans.

Human brucellosis is a zoonotic disease, that is, it is readily passed to humans from other species. Infection in humans is normally acquired either through consumption of contaminated dairy and meat products or by contact with infected animal secretions. Human beings are susceptible to *B. melitensis, B. suis, B. abortus* and *B. canis* in a decreasing order. Brucellosis in humans is characterized by undulant fever, headache, cold sweats and general malaise. The 45 disease can last from a few weeks to several years. If untreated, serious complications leading to death can occur.

Brucellosis among domestic livestock in North America has been largely controlled by using a combination of reliable and accurate diagnostic tests, removal of infected animals, and efficacious vaccines. However, this disease still exists among free-ranging wild life including feral swine (Sus scrofa). Infected wild life populations are the most likely source of transmission of brucellosis to humans, and for the possible reintroduction of this disease into domestic livestock. Feral swine populations are present in many regions of the world. Approximately 2 to 3 million feral swine are estimated to be present in the US alone, and feral swine populations in the southern portion of the US are known to be infected with B. suis.

Brucellosis among US domestic pig populations is currently controlled by depopulation procedures, a less than ideal strategy. Further, swine brucellosis is recognized as a major threat to domestic pig production in other parts of world. So far, no vaccine has been extensively used or clearly been proven useful against this disease in swine. 65 Therefore, it would be beneficial to have available a vaccine effective against brucellosis in feral and domestic swine.

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B. suis was the first pathogenic organism weaponized by the U.S. military during the 1950 s. Today it constitutes a potential bioterrorism threat that could be targeted against military personnel, civilians, or food supplies. Early diagnosis of brucellosis is problematic, and the treatment regiment is prolonged antibiotic therapy. However, antibiotic-resistant strains of Brucella can be generated easily, and if such strains are used in bio-warfare, use of antibiotics to control brucellosis may not be effective. The Centers for Disease Control and Prevention has listed Brucella as a category-B biothreat-agent. Currently, there is no licensed vaccine against brucellosis in humans. Due to its highly infectious nature and the increased likelihood of illegitimate use, it would be beneficial to have available a vaccine that protects humans against this pathogen.

Several animal vaccines against different strains of *Brucella* currently exist.

Cattle vaccine strain RB51: This is an attenuated (less capable of surviving in animals, and less capable of causing disease in animals), rough (incapable of producing the cell-surface antigen called O-side-chain) strain of *B. abortus*. Strain RB51 induces strong cell-mediated immune (CMI) responses and provides protection against brucellosis in bovine and several other animal species. It is the official vaccine approved by USDA to protect cattle against infection with *B. abortus*.

Although very effective in immunizing cattle against *B. abortus*, it is less effective against *B. melitensis* and *B. suis* infections, suggesting that strain RB51 would not be a suitable vaccine for humans, where *B. melitensis* and *B. suis* cause the most severe symptoms. The induction of O-side chain antibodies, in addition to strong CMI, appears to be important for protection against brucellosis in humans. Strain RB51 is rough, and therefore expresses only minimal amounts of O-side chain antigen and does not induce O-side chain antibodies.

In addition, strain RB51 was developed through natural selection procedures, and therefore, its genetic make up is not filly known. Further, strain RB51 is resistant to rifampicin, one of the very few antibiotics available for treatment of humans against brucellosis. Thus, if a human vaccinated with RB51 did become infected, (e.g. an immuno-compromised individual), it would not be possible to treat the infection with standard antibiotic therapy with rifampicin. For these reasons, strain RB51 is not considered a suitable candidate for use as a brucellosis vaccine for humans.

Cattle vaccine strain 19: This strain is able to induce protective immunity in cattle. However, although strain 19 (also known as S19) is of low virulence for cattle, vaccination of pregnant cows can still result in abortions. A less frequent adverse consequence of strain 19 vaccination is the development of an arthropathy associated with *Brucella* antigen-containing immune complexes (but not live organisms) in the affected joints.

Strain 19 is known to be pathogenic for human beings. In addition, this strain was isolated through laboratory selection procedures, and its genetic make up is not understood. Therefore, strain 19 is also not considered a suitable candidate for use as a brucellosis vaccine for humans.

Sheep/goat vaccine strain Rev1: Rev1 vaccine is a live, attenuated *B. melitensis* strain that stimulates protection against infection with *B. melitensis* in sheep and goats and also protects rams against infection with *B. ovis*. Depending on the dose administered during pregnancy, abortions will occur with variable frequency. In cattle, Rev1 gives better protection than strain 19. However, strain Rev1 is also not considered safe as a human vaccine because its genetic makeup is not known, it is pathogenic for human beings, and it is resistant to the antibiotic streptomycin.

### SUMMARY OF THE INVENTION

The invention provides compositions and methods for treating and preventing Brucellosis. The methods involve eliciting an immune response to pathogenic, virulent bacteria of the genus *Brucella* by administering a composition comprising attenuated, recombinant *Brucella* strains that exhibit a deficiency in carboxy-terminal protease (CtpA) activity. The compositions may be used as a vaccine in mammals including without limitation swine, reindeer, cattle and humans. In a preferred embodiment, the attenuated *Brucella* strain is a *Brucella suis* strain in which a portion of the CtpA gene has been deleted.

It is an object of this invention to provide an attenuated, recombinant *Brucella* strain with a deficiency in carboxylterminal processing protease (CtpA) activity. In one embodiment, the attenuated, recombinant *Brucella* strain is of the species *Brucella suis*. In some embodiments, the CtpA deficiency is caused by deletion of at least a portion of a gene encoding CtpA in the attenuated, recombinant *Brucella* strain. In one embodiment, the attenuated, recombinant *Brucella* strain. In one embodiment, the attenuated, recombinant *Brucella* strain is *Brucella abortus*, *Brucella suis*, *Brucella melitensis*, *Brucella neotomae*, *Brucella canis*, or *Brucella ovis*; and the mammal may be a human, swine, cattle or reindeer.

The invention also provides a method for eliciting an immune response to a Brucella species in a mammal, or treating or preventing Brucellosis in a mammal, including vaccinating a mammal against Brucellosis. The method comprises the step of administering to the mammal in a quantity sufficient to elicit an immune response, an attenu- 30 ated, recombinant Brucella strain with a deficiency in carboxyl-terminal processing protease (CtpA) activity. The attenuated, recombinant Brucella strain may be of the species Brucella suis, and the CtpA deficiency may be caused by deletion of at least a portion of a gene encoding CtpA in 35 the attenuated, recombinant Brucella strain. In one embodiment, the attenuated, recombinant Brucella strain is 1330∆ctpA. The *Brucella* species may be *Brucella abortus*, Brucella suis, Brucella melitensis, Brucella neotomae, Brucella canis, or Brucella ovis, and the mammal may be human, swine, cattle or reindeer.

The invention further provides a composition for eliciting an immune response to *Brucella* species in a mammal. The composition comprises an attenuated, recombinant *Brucella* strain with a deficiency in carboxyl-terminal processing protease (CtpA) activity and, a physiologically suitable carrier. The attenuated, recombinant *Brucella* strain may be of the species *Brucella suis*, and the CtpA deficiency may be caused by deletion of at least a portion of a gene encoding CtpA in the attenuated, recombinant *Brucella* strain. In one embodiment, the attenuated, recombinant *Brucella* strain is 50 1330ΔctpA. Further, the *Brucella* species may be *Brucella abortus*, *Brucella suis*, *Brucella melitensis*, *Brucella neotomae*, *Brucella canis*, or *Brucella ovis*, and the mammal may be humans, swine, cattle or reindeer.

The invention further provides a gene having a nucleotide sequence selected from SEQ ID NO: 1, SEQ ID NO: 3 and SEO ID NO: 4.

The invention also provides a method of detecting exposure of a mammal to *Brucella* species. The method comprises the steps of obtaining a biological sample from the mammal, and amplifying nucleic acid in the biological sample by polymerase chain reaction using primers specific for SEQ ID NO: 1 or SEQ ID NO: 3.

In another preferred embodiment, the invention provides an attenuated, recombinant *Brucella* strain with a deficiency in tail-specific protease (TspA) (a homologue of carboxyl-65 terminal processing protease) activity, and the deficiency is caused by a deletion in the TspA gene of *Brucella melitensis*.

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(TspA is a homologue of CtpA.) In one embodiment of the invention, and the attenuated, recombinant Brucella is  $16M\Delta tspA$ 

The invention further provides a method for eliciting an immune response to a *Brucella* species in a mammal, or for treating or preventing Brucellosis in a mammal, including vaccinating a mammal against Brucellosis. The method comprises the step of administering to the mammal in a quantity sufficient to elicit an immune response, an attenuated, recombinant *Brucella* strain with a deficiency in tail specific protease activity, in which the *Brucella* strain is *Brucella* melitensis, and the deficiency is caused by a deletion in TspA. In one embodiment of the method, the attenuated, recombinant *Brucella* strain is 16MAtspA, and the mammal is of a type selected from the group consisting of humans, sheep, goats, dogs, swine, cattle and reindeer.

The invention further provides a composition for eliciting an immune response to *Brucella* species in a mammal. The composition comprises an attenuated, recombinant *Brucella* strain with a deficiency in tail-specific protease (TspA) (a homologue of carboxyl-terminal processing protease) activity, in which the attenuated, recombinant *Brucella* strain is of the species *Brucella melitensis* and the is caused by a deletion in TspA. The composition further includes a physiologically suitable carrier. In one embodiment, the attenuated, recombinant *Brucella* strain is 16MðtspA.

The invention further provides a gene having a nucleotide sequence selected from SEQ ID NO: 7, SEQ ID NO: 9 and SEQ ID NO: 11.

The invention further provides a method of detecting exposure of a mammal to *Brucella* species. The method comprises the steps of 1) obtaining a biological sample from a mammal, and 2) amplifying nucleic acid in the biological sample by polymerase chain reaction using primers specific for SEQ ID NO: 7 or SEQ ID NO: 9.

#### BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1. Nucleotide (SEQ ID NO: 1) and amino acid (SEQ ID NO: 2) sequences of CtpA gene of *Brucella suis*.

FIG. 2. Sequence of the 471 nucleotides that were deleted from CtpA (SEQ ID NO: 3).

FIG. 3. Nucleotide sequence of  $1330\Delta \text{ctpA}$  (SEQ ID NO: 4).

FIG. 4. Physical map of the insert of pGEMΔctpAK suicide vector. Solid line indicates the length and location of the ctpA gene. Solid-unfilled arrows indicate the locations of primers used to PCR amplify a portion of the ctpA gene. Solid-fill block indicates the gene fragment amplified by PCR. Dotted rectangle indicates the region deleted by mutagenesis. The checked solid block indicates the inserted kan<sup>r</sup> gene.

FIGS. 5A and B. Growth of *B. suis* strains. Single colonies of strains 1330, 1330ΔctpA, and 1330ΔctpA[pBBctpA] were grown overnight in TSB for 48 hours. The cells were pelletted in two equal aliquots by centrifugation. One pellet was resuspended in 1 ml of regular LB broth and used to inoculate 25 ml regular LB broth in Klett side-arm flask to 12 to 16 Klett units. The other pellet was resuspended in salt-free LB media and used to inoculate 25 ml salt-free LB broth in Klett flask to 8 to 16 Klett units. All cultures were grown at 42° C. at 180 rpm. Klett readings were recorded every two hours in a Klett Sumerson calorimeter. FIG. 5A: Growth of strains in regular LB media. FIG. 5B: Growth of strains in salt-free LB media.

FIG. 6. Splenic clearance of *B. suis* strains in bi-weekly intervals. Groups of 25 mice each were intraperitoneally inoculated with 4.0-4.1 log<sub>10</sub> CFU of strains 1330 or

 $1330\Delta ctpA$  and the splenic CFU counts were determined 1, 3, 5, 7, or 9 weeks post-inoculation.

FIG. 7. ELISA detection of IgG1, IgG2a and total IgG antibodies in serum of mice vaccinated with strain 1330 $\Delta$ ctpA or inoculated with saline alone. Sera collected from eight mice of each group at 6 weeks post-vaccination were diluted 1/100 and assayed for the presence of specific antibodies. Sera collected from mice vaccinated with strain RB51 (obtained from A. Contreras, Virginia Tech) were used as a control. Results are shown as the mean of  $OD_{450}$  of the 10 color developed.

FIG. **8**. Cell morphology of strain 1330 grown in LB media with salt, determined by scanning electron microscopy. The cells possessed the native coccobacillus shape of *Brucella*. Additionally, the cells revealed the typical ultrastructure of Gram negative bacteria, namely, the outer membrane, periplasmic space, and cytoplasmic membrane. The magnification was  $\times 10,000$ .

FIG. 9. Cell morphology of strain 1330 grown in LB media without salt, determined by scanning electron microscopy. The cells possessed the native coccobacillus shape of *Brucella*, and typical ultrastructure of Gram negative bacteria. A significant difference cannot be seen in cell morphology of strain 1330, when salt is present or absent in growth media. The magnification was ×10,000.

FIG. 10. Cell morphology of strain 1330ΔctpA grown in LB media with salt, determined by scanning electron microscopy. The cells acquired a spherical shape instead of its native coccobacillus shape of *Brucella*. The cell diameter apparently increased slightly. Additionally, the cells partially lost the typical ultrastructure of Gram negative bacteria. The outer membrane apparently separated from some of the cells. However, the integrity of the rest of the cell was not changed. The magnification was ×10,000.

FIG. 11. Cell morphology of strain 1330 $\Delta$ ctpA grown in 35 LB media without salt, determined by scanning electron microscopy. The cells lost their native coccobacillus shape of *Brucella*. Additionally, the cells lost the typical ultrastructure of Gram negative bacteria. The outermembrane dissociated from cells. The integrity of the cell was changed. The 40 magnification was  $\times 10,000$ .

FIGS. **12**A and B. A, Nucleotide sequence encoding *Brucella melitensis* TspA (SEQ ID NO: 7); B, amino acid sequence of *Brucella melitensis* TspA (SEQ ID NO: 8).

FIG. 13. Protein sequence comparison for Brucella 45 melitensis TspA and other proteins. The deduced amino acid sequences of parts of the CtpA proteins and periplasmic proteins of A. tumefaciens (Agrobac) (GenBank accession no. NP\_355704.1; SEQ ID NO: 14), Bartonella quintana (Bartone) (GenBank accession no. Q44879; SEQ ID NO: 50 15), Bradyrhizobium japonicum (Bradyrh) (GenBank accession no. NP\_771462.1; SEQ ID NO: 16), Brucella melitensis (Brucell) (GenBank accession no.NP\_539132.1; SEQ ID NO: 17), Mesorhizobium loti (Mesorhi) (GenBank accession no NP\_104979.1; SEQ ID NO: 18), Rhodopseudomonas palustris (Rhodops) (GenBank accession no. ZP\_00009772.1; SEQ ID NO: 19), and Sinorhizobium meliloti (Sinorhi) (GenBank accession no. NP\_387272.1; SEQ ID NO: 20) were compared. The ClustalV (PAM250) program of DNASTAR was used to align the sequences. The numbers on the left are the positions of the amino acid 60 residues in the proteins. The shaded amino acids are identical in all proteins. Amino acids shown to be essential for enzymatic activity in Synechocystis sp. Strain PCC 6803 and B. burgdorferi strain B31 are indicated by solid circles.

FIGS. **14**A, B and C.A, Nucleotide sequence encoding 65 *Brucella melitensis* deletion mutant 16MΔtspA; (SEQ ID NO: 9) B, amino acid sequence of *Brucella melitensis* 

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deletion mutant  $16\text{M}\Delta \text{tspA}$  (SEQ ID NO: 10); C, nucleotide sequence that was deleted from the tspA gene (SEQ ID NO: 11)

FIGS. **15**A and B. Growth of *B. melitensis* strains 16M (♠), 16MΔtspA (♠), and 16MΔtspA[ctpA<sup>+</sup>] (♠). All cultures were grown at 37° C. at 200 rpm. Changes in cell density were recorded every two hours in a Klett-Summerson colorimeter. A, Growth of strains in LB media. B, Growth of strains in salt-free LB media.

FIGS. **16**A and B. The cell morphology of *B. melitensis* strains as observed by electron microscopy. The strain 16M 16 A) displayed native coccobacillus shape of *Brucella*, whereas, those from the strain 16MΔtspA (16 B) acquired a spherical shape with some cell with increased cell diameter. The 1-micron scale bar is inserted in each figure. The magnification is ×10,000.

# DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS OF THE INVENTION

The present invention provides compositions and methods for preventing or treating Brucellosis in mammals. The compositions elicit an immune response against virulent, pathogenic Brucella species, and thus may be used as vaccines. The compositions comprise at least one attenuated recombinant Brucella bacterial strain in which there is a deficiency in carboxyl-terminal protease (CtpA) activity. CtpA activity may be encoded by a ctpA gene (as in Brucella suis). Alternatively, CtpA activity may be encoded by a homologue with carboxyl-terminal protease activity designated as tail-specific protease (such as the tspA gene of Brucella melitensis). By "deficiency" we mean that CtpA activity is partially or totally absent in the bacterium. For example, the gene encoding CtpA activity may be partially or totally deleted from the Brucella bacteria, e.g. by genetic engineering techniques as described herein. Due to the use of such genetic engineering techniques, the genetic makeup of the attenuated strain is fully known, an advantage for a vaccine composition. Alternatively, the gene encoding CtpA (e.g. CtpA or TspA) may be altered in some other manner that inactivates the gene, or greatly reduces its activity, e.g. by the introduction of mutations within the gene by genetic engineering, thereby reducing or eliminating the function of the gene and/or of the CtpA protein encoded by the gene, by preventing transcription or translation of the gene, etc. In any case, whether by deletion or by some other mutation, the result is that CtpA activity within the Brucella bacteria is modified so as to be non-existent or greatly reduced. By "greatly reduced" we mean that the modification results in a reduction in activity (compared to CtpA activity in wild type cells) of at least about 50-100%, preferably about 75-100%, and most preferably from about 90-100%. This reduction in CtpA activity may be due, for example, to mutations that are introduced into the CtpA gene to cause a severe reduction in the amount of CtpA that is produced in the cell, or to cause the form of CtpA that is produced by the mutated gene to be greatly reduced in activity or to be non-active. Those of skill in the art are well acquainted with procedures for assaying the level of activity of CtpA in Brucella bacteria, and methods for the comparison of levels of enzyme activity between wild type and attenuated bacteria are also known (see, for example, methods described in the Examples section herein). Enzyme activity may be measured directly, or may be inferred by measurement of some other observable trait, e.g. growth rate, sensitivity to temperature, medium conditions, etc., or by a combination of both. Further, genetic changes in the bacteria may be detected/ confirmed by techniques familiar to those of skill in genetic

analyses, as also described herein (e.g. polymerase chain reaction (PCR) of bacterial genetic material with suitable primers).

Brucella strains that display such a deficiency in CtpA activity are "attenuated", that is the bacteria that are used in 5 the composition are living and able to reproduce but compared with the infectious strains, they are less capable of surviving in animal or human hosts, and incapable or less capable of causing disease in hosts. The level of attenuation of a strain is determined using mouse experimental models. In this work, mice are injected with the strains, and at different time intervals (1, 3, 5, 7 and 9 weeks after injection), injected mice are sacrificed, their spleens are isolated and crushed, spleen content is plated on growth media plates, and incubated for 3-5 days at 37° C. at the presence of carbon-dioxide. Each cell of *Brucella* present in a spleen is expected to form a colony (colony forming unit or CFU) on plates after 3-5 days of incubation. If the injected strain is attenuated, its presence in the spleen never increases, but gradually declines with time and eventually disappears completely from the spleen. If the presence of a strain in 20 spleen declines to about 100-600 CFU in 5-7 weeks, and disappears completely in 6-9 weeks after injection, it can be considered sufficiently attenuated or safe to be a vaccine. On the contrary, if a strain is infectious (virulent), its presence in spleens increases to about 1,000,000 CFU one week after 25 injection and remains at a constant level of approximately 3000 CFU for more than 9 weeks after injection. Those of skill in the art are acquainted with procedures for growing attenuated Brucella species for the production of compositions for use as vaccines, for example, those that are outlined 30 in the Examples section that is included herein.

The Brucella species or strain that is modified for use in the practice of the present invention (i.e. the species or strain which is genetically manipulated to produce or derive the recombinant, attenuated Brucella bacteria) may be any suitable *Brucella* species or strain. Examples include but are not limited to Brucella abortus, Brucella canis, Brucella melitensis, Brucella neotomae and Brucella ovis, Brucella suis, and various strains thereof. In a preferred embodiment of the invention, the attenuated Brucella bacteria are derived from  $Brucella\ suis$ . In addition, in one embodiment of the  $^{40}$ invention, the recombinant, attenuated Brucella suis has a deletion in the CtpA gene. In a preferred embodiment of the invention, the attenuated Brucella suis strain is recombinant strain1330ΔctpA. 1330ΔctpA is derived from Brucella suis and has a 471 base pair deletion in the ctpA gene. The 45 nucleotide sequence of the ctpA gene (SEQ ID NO: 1) and the corresponding amino acid sequence (SEQ ID NO: 2) are given in FIG. 1. FIG. 2 depicts the 471 basepairs that are deleted and FIG. 3 (SEQ ID NO: 3) shows the nucleotide sequence of the CtpA gene after deletion of the 471 base  $_{50}$ pairs (SEQ ID NO: 4). However, those of skill in the art will recognize that it is not necessary to delete precisely this 471 basepair segment of the gene in order to generate an attenuated Brucella strain for use in the practice of the present invention. For example, more of the gene (e.g. up to 100% of the nucleotide sequences encoding the CtpA gene) may be deleted. Thus, in one embodiment, at least the indicated 471 base pair region is deleted. Alternatively, somewhat less extensive deletions may be employed, or deletions in other regions of the gene may be made, or deletions that overlap the 471 base pair region may be made, so long as the 60 resulting Brucella bacteria are attenuated and can be used to elicit an immune response against at least one virulent Brucella species, and preferably provide protection against infection by a virulent Brucella species, in at least one mammal of interest.

In another preferred embodiment, the *Brucella* species is *Brucella melitensis* and the gene that encodes carboxy-

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terminal processing activity (also known as tail-specific protease activity) is the tspA gene, a homologue of the ctpA gene in *Brucella suis*. A preferred example of this embodiment is the case in which 471-base pair region is deleted from the tspA gene, to form the attenuated *Brucella melitensis* strain 16MΔtspA. The nucleotidesequence of TspA from *Brucella melitensis* (SEQ ID NO:7) is given in FIG. 12A, and the amino acid sequence (SEQ ID NO: 8) is given in FIG. 12B. The deleted forms of TspA present in the attenuated *Brucella melitensis* 16MΔtspA are depicted in FIG. 13A (nucleotide, SEQ ID NO: 9) and B (amino acid, SEQ ID NO: 10), as is the deleted nucleotide sequence (SEQ ID NO: 11, FIG. 13C).

In some embodiments, a suitable section (or all) of the ctpA gene is deleted from the *Brucella* chromosome. Alternatively, a suitable section of the ctpA gene may be replaced by a different nucleotide sequence, e.g. by a sequence which facilitates selection of deletions mutants such as an antibiotic resistance gene, or a non-antibiotic selection marker such as sacB or leuB. Techniques for performing such replacements are known, and include the technique of allelic exchange, as utilized and described herein.

In the practice of this invention, it is desirable to either delete or otherwise disable (gene replacement, etc.) the CtpA sequence in a *Brucella* strain such that antibody titer to the strain can be raised but where the attenuated strain is deficient in CtpA activity.

The invention further provides the genetic sequence of the ctpA gene of *Brucella suis*, as depicted in FIG. 1, as well as some exemplary sequences of a gene with a deletion. These sequences might be used for transfecting an organism to incorporate a gene encoding CtpA, as a probe to identify organisms which harbor genes encoding CtpA, as well as to identify related genes in other species based on homology and other factors, as well as for other applications. The ctpA gene may also find application for use in combination with other genes of interest where carboxyl terminal protease activity is desired. Furthermore, the transfected proteins from these sequences may have similar applications.

Those of skill in the art will recognize that many variants of the sequence may exist or be constructed which would also function in the practice of the present invention. For example, with respect to amino acid sequences, variants may exist or be constructed which display: conservative amino acid substitutions; non-conservative amino acid substitutions; truncation by, for example, deletion of amino acids at the amino or carboxy terminus, or internally within the molecule; or by addition of amino acids at the amino or carboxy terminus, or internally within the molecule (e.g. the addition of a histidine tag for purposes of facilitating protein isolation, the substitution of residues to alter solubility properties, the replacement of residues which comprise protease cleavage sites to eliminate cleavage and increase stability, the addition or elimination of glycosylation sites, and the like, or for any other reason). Such variants may be naturally occurring (e.g. as a result of natural variations between species or between individuals); or they may be purposefully introduced (e.g. in a laboratory setting using genetic engineering techniques). All such variants of the sequences disclosed herein are intended to be encompassed by the teaching of the present invention, provided the sequence displays sufficient identity to the described sequences. Preferably, identity will be in the range of about 50 to 100%, and more preferably in the range of about 75 to 100%, and most preferably in the range of about 80 to 100% of the disclosed sequences. The identity is with reference to the portion of the amino acid sequence that corresponds to the original antigen sequence, i.e. not including additional elements that might be added, such as those described below for chimeric antigens. Further, all sequences which hybrid-

ize to the depicted sequence under stringent hybridization conditions are also encompassed.

The present invention provides compositions for use in eliciting an immune response, and which may be utilized as a vaccine against Brucellosis. By "eliciting an immune 5 response" we mean that the composition stimulates synthesis of specific antibodies against the attenuated, recombinant Brucella strain at a titer of from about 1 to about  $5\times10^6$  or greater. In some embodiments, the titer is at least in the range of about 100 to about 1000 (or more), as measured by techniques that are known to those of skill in the art, for example, by <sup>3</sup>H thymidine incorporation or by Enzyme Linked Immunosorbent Assay (ELISA). In a preferred embodiment, the titer is measured by ELISA and the antibody titer is about  $3.4 \times 10^2$  to about  $4.3 \times 10^2$ . Further, the antibodies that are produced cross-react with at least one other pathogenic, virulent Brucella species or strain, against which it is desired to raise an immune response.

The compositions of the present invention include substantially purified attenuated, recombinant Brucella bacteria with a deficiency in CtpA activity, and a pharmacologically 20 suitable carrier. The preparation of such compositions for use as vaccines is well known to those of skill in the art. Typically, such compositions are prepared either as liquid solutions or suspensions, however solid forms such as tablets, pills, powders and the like are also contemplated. 25 Solid forms suitable for solution in, or suspension in, liquids prior to administration may also be prepared. The preparation may also be emulsified. The active ingredients may be mixed with excipients that are pharmaceutically acceptable and compatible with the active ingredients. Suitable excipients are, for example, water, saline, dextrose, glycerol, ethanol and the like, or combinations thereof. In addition, the composition may contain minor amounts of auxiliary substances such as wetting or emulsifying agents, pH buffering agents, and the like. In addition, the composition may contain other adjuvants. If it is desired to administer an oral form of the composition, various thickeners, flavorings, diluents, emulsifiers, dispersing aids or binders and the like may be added. The composition of the present invention may contain any such additional ingredients so as to provide the composition in a form suitable for administration. The 40 final amount of active ingredient (i.e. the attenuated Brucella strain) in the formulations may vary. However, in general, the amount will be from about 1-99% of the total composition. The vaccine preparations of the present invention may further comprise an adjuvant, suitable examples of 45 which include but are not limited to Seppic, Quil A, Alhydrogel, etc.

The present invention provides not only compositions, but also methods for their use to elicit an immune response. By "elicit an immune response", we mean that administration of 50 the composition causes the synthesis of specific antibodies at a titer in the range of from about 1 to about  $1 \times 10^6$  or greater. Preferably, the titer is from about 10,000 to about  $1\times10^6$  or more, and most preferably, the titer is greater than  $1\times10^6$  as measured, e.g. by <sup>3</sup>H thymidine incorporation. The methods involve administering a composition comprising attenuated Brucella strains with a deficiency in CtpA activity in a pharmacologically acceptable carrier to a mammal. The vaccine preparations of the present invention may be administered by any of the many suitable means which are well known to those of skill in the art, including but not limited 60 to by injection (e.g. subcutaneous or intramuscular), orally, intranasally, by ingestion of a food product containing the antigen, etc. In preferred embodiments, the mode of administration is subcutaneous or intramuscular.

The methods of the present invention are directed to <sup>65</sup> eliciting an immune response in a mammal. In some embodiments, the mammal is an animal such as sheep,

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goats, dogs, swine, reindeer, and cattle (either domestic or feral). In another embodiment of the invention, the mammal is a human. Where the disease of Brucellosis is involved, those of skill in the art will recognize that many strains of Brucella infect more than one species of mammal. Thus, a composition for vaccinating mammals against Brucella need not be specific for the species being vaccinated so much as efficacious against particular Brucella strains, since a strain can infect several species. The compositions of the present invention may be used to vaccinate mammals of any species, so long as they are infected with or are at risk for being infected with a strain or species of Brucella to which the immune response elicited by the compositions is relevant, i.e. in which the immune response elicited by the compositions is effective against treating or preventing disease symptoms that would otherwise be caused by the Brucella strain/species. Those of skill in the art are well versed in procedures for determining the efficacy of a composition to elicit an immune response to Brucella bacteria, for example, those that are discussed in the Examples section herein. In general, in order for a composition to be considered effective as a vaccine, the following criteria are used:

- (1) Attenuated (as described above)
- (2) Induce immune responses (as described above)
- (3) Induce protection in animals against infection (challenge) with the pathogenic (virulent) strain of Brucellain determining this aspect, groups of mice are injected (vaccinated) intraperitoneally with either saline, or the vaccine strain (i.e. strain 1330ΔctpA). About eight weeks after vaccination, all mice are injected (challenged) intraperitoneally with pathogenic (virulent) strains of Brucella (i.e., B. abortus strain 2308, B. melitensis strain 16M, or B. suis strain 1330). Two weeks after challenge, mice are sacrificed, and the presence of pathogenic Brucella in spleens will be determined, as described previously. In mice injected with saline, immune responses are not developed and therefore, the challenged virulent Brucella strains (2308, 16M or 1330) are expected to retain and multiply. Therefore, from spleens of those mice (injected with saline) 100,000 to 1000,000 or more CFU of challenged Brucella can be recovered. On the contrary, of those mice injected with the vaccine strain, immune responses are expected to develop, and as a result, most of the challenged Brucella are expected to be lysed (destroyed). From spleens of these mice (injected with vaccine) 0 to 600 CFU of challenged Brucella can be recovered. In a preferred embodiment of the invention, the composition is active in eliciting an immune response against Brucella species that include but are not limited to Brucella abortus, Brucella suis, Brucella melitensis, and Brucella ovis.

In addition, the compositions of the present invention may be used either prophylactically to prevent a mammal from contracting Brucellosis, or after the fact to treat a known (or suspected) infection in order to ameliorate symptoms of the disease.

The invention also provides two methods of detecting *Brucella* infection. In particular, the method is useful for differentiating infectious, virulent field strains of *Brucella* from the attenuated recombinant strains of the present invention. The first method involves obtaining a suitable biological sample from a mammal (e.g. mice, swine or cattle), and carrying out polymerase chain reaction (PCR) on the sample using primers that specifically amplify the ctpA gene. By amplifying the ctpA gene of FIG. 1, *Brucella* infection or previous *Brucella* exposure (e.g. by vaccination or otherwise) may be identified. If the PCR using the biological sample amplifies a 1408 basepair size fragment,

it can be concluded that the respective animal has been infected with a field isolate of Brucella. If the PCR amplifies a 2189 basepair size fragment, it is an indication that the respective animal has been vaccinated with the invented strain 1330 $\Delta$ ctpA. The second method involves obtaining a suitable biological sample from a mammal (e.g. mouse, swine or cattle), isolating the bacterium from the sample, and growing it in growth media with or without salt. If the isolated strain grows in both media (with salt or without salt), it is an indication that the respective animal has been infected with a field isolate of Brucella. If the isolated bacterium grows in media that contains salt but does not grow in media that does not contain salt, it indicates that the respective animal has been vaccinated with the invented strain 1330 $\Delta$ ctpA.

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#### **EXAMPLES**

#### Example 1

Animal brucellosis is a disease affecting various domestic 20 and wild life species, resulting from infection with bacteria belonging to the genus Brucella (Corbel and Brinley Morgan, 1984). Brucellosis is a zoonotic disease and human infection is normally acquired either through consumption of contaminated dairy and meat products or by contact with 25 infected animal secretions (Acha and Szyfres, 1980). Brucella species are facultative intracellular pathogens that enter the host via mucosal surfaces and are able to survive inside macrophages. The primary strategy for survival in macrophages appears to be inhibition of phagosome-lysosome fusion (Arenas et al., 2000; Baldwin and Winter, 1994; Naroeni et al., 2001). Localization and survival within autophagosome-like compartments associated with the rough endoplasmic reticulum has also been demonstrated in placental trophoblasts and other non-professional phagocytes (Anderson et al., 1986; Pizarro-Cerda et al., 1998). Molecular characterization of this survival process is important because it would provide additional guidance for the development of measures for prevention and control of Brucella and perhaps other intracellular pathogens. As the result of annotating the B. suis genome (Paulsen et al., 40 2002), putative virulence genes in B. suis are being identified by looking for virulence homologs that have been reported in other pathogens.

It is well known that many proteins destined for extracytoplasmic locations are initially synthesized as precursor 45 forms and processed into mature forms by proteolytic cleavage to remove short peptide sequences, either near the amino terminus or near the carboxyl terminus of such proteins. The endoproteases responsible for cleaving of amino-terminal peptides are called amino-terminal processing proteases and 50 have been identified and studied in a number of systems. During recent years, a relatively new class of endoproteases with carboxyl-terminal processing activities has been described in various bacteria and organellar systems including cyanobacteria, chloroplasts, and E. coli (Keiler and Sauer, 1998; Pakrasi, 1998; Satoh, 1998; Silber et al., 1992; Keiler, et al., 1996). These carboxyl-terminal proteases (Ctp) from cyanobacteria, E. coli and green plants share significant sequence similarities (Inagaki et al., 1996; Oelmüller et al., 1996). However, none of them exhibits sequence homology with other protease classes with well-defined mechanisms of action. Ctps are serine proteases that utilize a Ser/Lys catalytic dyad instead of the well-known Ser/His/ Asp catalytic triad (Paetzel and Dalbey, 1997).

The enzymes involved in synthesis of the bacterial cell wall are named as penicillin binding proteins (PBPs). This 65 name has been given to these enzymes because b-Lactam antibiotics, including penicillin, bind covalently and irre-

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versibly to these enzymes and inhibit the synthesis of the peptidoglycan layer (Waxman and Strominger, 1983). In E. coli, eight PBPs have been identified. Among them, PBP 3 is believed to be involved in polar cap murein synthesis/cell division (Yousif et al., 1985). In E. coli, the bulk of the PBP 3 molecule, except for the N-terminal membrane anchor region, protrudes into the periplasmic space, where it acts on murein (Bowler and Spratt, 1989). Amino acid sequence analysis of precursor and mature forms of PBP 3 (Nagasawa et al., 1989) revealed that cleavage of eleven C-terminal residues is responsible for the maturation of PBP 3 protein. The C-terminal protease Prc has been identified as responsible for cleavage of the C-terminal 11 amino acid residues from the PBP 3 precursor. The Prc protein resides on the outer side of the cytoplasmic membrane (Hara et al., 1991). The E. coli mutant JE7304, developed by Hara et al., (1989) by deleting the prc gene encoding Prc protein was defective in the C-terminal processing of PBP 3. This mutant showed thermo-sensitive growth on a salt-free L-agar plate, suggesting that the prc gene is involved in some essential cellular process, which may or may not be related to the cell division function of PBP 3 (Hara et al., 1991). The prc function thus seemed to be involved in protection of the cell from thermal and osmotic stresses. Loss of Prc function also resulted in leakage of periplasmic proteins including RNase I and alkaline phosphatase (Hara et al., 1991). The leaky phenotype of the prc mutant has been attributed to the impairment of the structural integrity of the outer membrane, which could lead to sensitivity to osmotic stress (Hara et al., 1991).

The ctpA gene is 1274-bp long and is located between 1768433 and 1769707-bp on chromosome I of the *B. suis* genome. The predicted molecular mass of CtpA is 45.2-kDa. The protein encoded by this gene shares 31% homology with Prc protein of *E. coli* and up to 77% homology with the Ctps of other bacteria. Based on this homology, it was hypothesized that the protein encoded by ctpA was a C-terminal protease that could play a significant role in determining the virulence of *B. suis*. It is herein reported that a *B. suis* strain with a defective ctpA gene exhibits salt-sensitive growth exactly as the Prc-deficient *E. coli* did. In addition, this strain produces smaller colonies on enriched agar plates, and exhibits slow growth in enriched growth media and reduced persistence in mice and mouse macrophages.

#### Materials and Methods

Bacterial strains, plasmids, and reagents. B. abortus strain 2308, B. melitensis strain 16M, and B. suis strains 1330 and VTRS1 were obtained from our culture collection. E. coli strain Top10 (Invitrogen Life Technologies, Carlsbad, Calif.) was used for producing plasmid constructs. E. coli Prc mutant strain JE7929 was a gift. E. coli were grown in Luria Bertani (LB) broth or on LB agar (Difco Laboratories, Sparks, Md.). Brucellae were grown in LB broth with or without sodium chloride at 30, 37 or 42° C. to determine whether growth was osmo-sensitive and/or thermo-sensitive. For all other assays, Brucellae were grown either in trypticase soy broth (TSB) or on trypticase soy agar (TSA) plates (Difco) at 37° C. as previously described (Schurig et al., 1991). The plasmids used in this study are listed in Table 1. Bacteria containing plasmids were grown in the presence of ampicillin or kanamycin at 100-µg/ml concentration (Table 1).

All experiments with live Brucellae were performed in a Biosafety Level 3 facility at the Infectious Disease Unit of the Virginia-Maryland Regional College of Veterinary Medicine.

TABLE 1

Description of the plasmids and bacterial strains used in this study				
Name	Description	Source or reference		
Plasmids				
pCR2.1 pCRctpA	TA cloning vector, 3.9-kb; Amp <sup>r</sup> pCR2.1 with 1.4-kb insert containing	Invitrogen This study		
pGEM-3Z pGMEctpA	the <i>B. suis</i> ctpA gene; Amp <sup>r</sup> Cloning vector, 2.74-kb, Amp <sup>r</sup> pGEM-3Z with a 1.4-kb insert containing the <i>B. suis</i> ctpA gene	Promega This study		
pUC4K pGEMctpAK	from pCRctpA; Amp <sup>r</sup> Cloning vector, 3.9-kb, Kan <sup>r</sup> , Amp <sup>r</sup> PGEMctp with 0.5-kb BcII fragment deleted and blunt ended and a 1.3-kb SaII-cut and blunt-ended Kan <sup>r</sup>	This study		
pBBR4MCS pBBctpA	cassette from pUC4K ligated, Kan <sup>r</sup> , Amp <sup>r</sup> Broad-host range vector; Cm <sup>r</sup> pBBR4MCS with a 1.4-kb insert containing the <i>B. suis</i> ctpA gene from pCRctpA; Amp <sup>r</sup>	Kovach et al., 1994 This study		
Bacteria				
Escherichia coli				
Top 10	F-mcrAΔ(mrr-hsdRMS-mcrBC) Φ80lacZΔM15ΔlacX74deoR recA1araD139Δ(ara-leu)7697 galUgalKrpsL (StrR) endA1 nupG	Invitrogen		
JE7929 B. abortus 2308 B. melitensis 16M B. suis 1330 1330ActpA 1330ActpA[pBBctpA]	Prc mutant Wild-type, smooth strain Wild-type, smooth strain Wild-type, smooth strain ctpA deleted mutant of 1330, Kan <sup>r</sup> Strain 1330 containing pBBctpA, Kan <sup>r</sup> , Amp <sup>r</sup>	Fraipont et al., 1994 G. G. Schurig G. G. Schurig G. G. Schurig This study This study		
VTRS1	wboA deletion mutant of B. suis	Winter (1996)		

Recombinant DNA methods. Genomic DNA was isolated from *B. suis* strain 1330 using a Qiagen Blood and tissue DNA kit (Qiagen Inc., Valencia, Calif.). Plasmid DNA was isolated using plasmid mini or midi prep purification kits (Qiagen). Restriction digests, Klenow reactions, and ligations of DNA were performed as described elsewhere (Sambrook et al., 2001). Restriction enzymes, Klenow fragment and T4 DNA ligase enzyme were purchased from Promega Corporation (Madison, Wis.). Ligated plasmid DNA was transferred to *E. coli* Top10 cells by heat shock transformation, as per manufacturer's guidelines (Invitrogen). Purified plasmid DNA was electroporated into *B. suis* with a BTX ECM-600 electroporator (BTX, San Diego, Calif.), as described previously (McQuiston et al., 1995).

DNA sequence analysis. The nucleotide sequence of ctpA gene was analyzed with DNASTAR software (DNASTAR, Inc., Madison, Wis.). Sequence similarity searches of the EMBL/GenBank/DDBJ databases were performed using BLAST software (Altschul et al., 1990) at the National Center for Biotechnology Information (Bethesda, Md.).

Mutation of the *B. suis* ctpA gene by allelic exchange. A 1408-bp region including a major portion of the ctpA gene was amplified via PCR using the genomic DNA of *B. suis* (FIG. 4). A primer pair consisting of a forward primer (5 ' GGGGTACCGTGGTGGACTGA 3 ') (SEQ ID NO: 5) and a reverse primer (5 ' GGCTGCAGTC-CCGCGTTTTTGTCTT 3 ') (SEQ ID NO: 6) (Ransom Hill Bioscience, Inc., Ramona, Calif) were designed based on the nucleotide sequence (GenBank accession no. NC\_004310). The *B. suis* genomic sequence 89 to 78-bp upstream from 65 ATG starting codon of ctpA gene was used to design the forward primer, whereas the sequence 14 to 37-bp down-

stream from the stop codon of ctpA was used to design the reverse primer (FIG. 4). A restriction site was engineered into each primer (KpnI in the forward primer, and PstI in the reverse primer, shown in bold case in the primer sequences). PCR amplification was performed in an Omni Gene thermocycler (Hybaid, Franklin, Mass.) at 95° C. for 5 min, followed by 35 cycles that each included 1 min of denaturation at 95° C., 1 min of annealing at 59.7° C., and 3 min of extension at 72° C. The amplified gene fragment was cloned into the pCR2.1 vector of the TA cloning system (Invitrogen) to produce plasmid pCRctpA. Competent E. coli Top10 cells (Invitrogen) were transformed with the ligation mixture, and the colonies carrying the recombinant plasmid were picked from TSA plates containing ampicillin (100 µg/ml), as per the manufacturer's guidelines. From this plasmid, the ctpA gene was isolated by KpnI and PstI digestion and cloned into the same sites of plasmid pGEM-3Z (Promega). The resulting 4.2-kb plasmid was designated as pGEMctpA. Competent E. coli Top10 cells (Invitrogen) were transformed with the ligation mixture, and the colonies carrying the recombinant plasmid were picked from TSA plates containing ampicillin (100 μg/ml). The suicide vector pGEMctpAK was constructed as follows: the plasmid pGEMctpA was digested with BclI to delete a 471-bp region from ctpA gene. The BclI sites on the 3.7-kb plasmid were filled in by reaction with Klenow enzyme and ligated to the 1.3-kb SalI fragment of pUC4K (also blunt ended) containing the Tn903 npt gene (Ried and Colmer, 1987), which confers kanamycin resistance (Kan<sup>R</sup>) to B. suis. The resulting suicide vector was designated pGEMctpAK. Competent E. coli Top10 cells (Invitrogen) were transformed with the ligation mixture, and

the colonies carrying the recombinant plasmid were picked from TSA plates containing kanamycin (100 µg/ml).

One microgram of pGEMctpAK was used to electroporate B. suis strain 1330; several colonies of strain 1330 were obtained from a TSA plate containing kanamycin (100 <sup>5</sup> μg/ml). These colonies were streaked on TSA plate containing ampicillin (100 μg/ml) to determine if a single- or double-crossover event had occurred. Three of the colonies did not grow on ampicillin containing plates suggesting that a double cross-over event had occurred, whereas the rest of 10 the colonies grew on ampicillin containing plates suggesting that a single cross-over event had occurred. PCR with the primers used for amplifying the ctpA gene (as described above) confirmed that a double-crossover event had taken place in all three transformants. Cells were harvested from 15 B. suis strains, boiled for 30 min, and centrifuged for 15 min. The supernatant was used for PCR using the forward and the reverse primers (FIG. 4). Strain 1330 amplified a 1408-bp fragment whereas strain 1330ΔctpA amplified a 2189-bp fragment. (data not shown). One of these strains was chosen 20 for further analyses and designated  $1330\Delta ctpA$ .

Complementation of ctpA gene activity in mutant 1330ΔctpA. The 1.4-kb DNA fragment containing *B. suis* ctpA gene was isolated by SacI and XbaI digestion of plasmid pCRctpA and was cloned into same sites of broadhost range vector pBBR4MCS (Kovach et al., 1994). The resulting plasmid was designated as pBBctpA. One microgram of pBBctpA was used to electroporate *B. suis* strain 1330ΔctpA; several colonies of strain 1330ΔctpA were picked from a TSA plate containing ampicillin (100 μg/ml). Six of the colonies were tested for growth in salt-free LB media (details below). One of the colonies that grew well in this media was chosen for further analyses and designated as 1330ΔctpA[pBBctpA].

Complementation of prc gene activity in Prc-deficient *E. coli*. One microgram of pBBctpA was used to electroporate the Prc mutant *E. coli* strain JE7929; several colonies of strain JE7929 were picked from a TSA plate containing ampicillin (100 µg/ml). Ten of the colonies were tested for growth in salt-free LB media (details to follow).

Growth rates of *B. suis* strains in regular or salt-free media at different temperatures. Salt-free LB media was prepared by mixing bactotryptone and yeast in water per manufacturer's instruction (Difco), but omitting sodium chloride. Single colonies of strains 1330, 1330ΔctpA, and 1330ΔctpA [pBBctpA] were grown at 37° C. for 24 hours to stationary phase in 10 ml of TSB. The cells were pelleted in two equal aliquots by centrifugation. One pellet was resuspended in 1 ml of regular LB broth and used to inoculate 25 ml of regular LB broth in a Klett side-arm flask to 12 to 16 Klett units. The other pellet was resuspended in salt-free LB media and used to inoculate 25 ml of salt-free LB broth in a Klett flask to 8 to 16 Klett units. Cultures were grown at 30, 37, or 42° C. at 180 rpm; Klett readings were recorded every two hours in a Klett Sumerson photometer.

Acid precipitation and denaturing gel electrophoresis of secreted proteins. Strains 1330, 1330ΔctpA and 1330ΔctpA [pBBctpA] were grown in 25 ml LB broth to stationary phase (to 339, 179 and 288 Klett units, respectively). The culture was centrifuged at 2000×G for 15 min, and the cell free culture medium was collected. Trichloroacetic acid (TCA) was added to the medium at 5% of final volume and incubated at 4° C. overnight. The acidified medium was centrifuged at 10000×g for 15 min to collect the protein precipitate. The insoluble material was resuspended in Laemmli sample buffer (Sigma Chemical Co., St. Louis, Mo.), boiled for 20 min and electrophoresed on 10% SDS/ 65 PAGE gels according to standard procedures (Laemmli, 1970). Gels containing the separated proteins were either

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stained with Coomassie brilliant blue G (Sigma Chemical Co.) or used for Western blot analysis.

Western blotting. Western blotting was performed as previously described (Vemulapalli et al., 1998). Briefly, proteins separated by SDS-PAGE were transferred to a nitrocellulose membrane by using a Trans-blot semidry system (Bio-Rad Laboratories, Hercules, Calif.). The membranes were blocked with a solution of 1.5% non-fat milk powder plus 1.5% bovine serum albumin. For analysis of TCA insoluble proteins, the membranes were incubated with goat anti-heat killed *B. abortus* polyclonal serum (Goat-48) for 24 hours and subsequently developed with rabbit antigoat IgG (whole molecule) conjugated with horseradish peroxidase (Sigma Chemical Co).

Phenotypic characterization of *B. suis* strains. Recombinant colonies were analyzed for their rough or smooth phenotype by crystal violet staining as described previously (Alton, et al., 1975; White and Wilson, 1951). Briefly, crystal violet staining solution was added to colonies grown on TSA plates. After two minutes, the solution was poured off and the colonies were observed to determine if they retained the stain (rough phenotype) or not (smooth phenotype).

Cell morphology *B. suis* strains. The wild type and the CtpA-deficient *B. suis* strains were grown in media with or without salt, and the cells were used for a Gram-staining. The stained specimens were observed using a light microscope at 100× magnification.

Preparation of *B. suis* inoculum stocks. TSA plates were inoculated with single colonies of *B. suis* strains. After four days of incubation at 37° C. in CO<sub>2</sub>, the cells were scooped/harvested from plates, washed with PBS, resuspended in 20% glycerol, and saved at -80° C. The number of viable cells was determined after culturing dilutions of the cell suspensions on TSB.

Persistence of recombinant B. suis strains in J774 macrophages. J774 macrophage cells were seeded at a density of 5×10<sup>5</sup>/ml in Dulbecco's modified essential medium (DMEM) (Sigma-Aldrich) into 24-well tissue culture dishes and cultured until confluent. The tissue culture medium was removed, 200  $\mu$ l (10<sup>8</sup> cells) of the bacterial suspension in PBS was added, and the cells were incubated at 37° C. for four hours. The suspension above the cell monolayer was removed, and the cells were washed three times with PBS. One milliliter of DMEM containing 25 µg of gentamicin/ml was added, and the cells were incubated for 48 hours at 37° C. At various time points (0, 4, 24 and 48 hours of incubation), the growth medium was removed, the cells were washed with PBS, and 500 µl of 0.25% sodium deoxycholate was added to the cells, which were lysed by aspiration. After 5 minutes the lysate was diluted in PBS, and the number of viable cells was determined after growth at 37° C. for 72 h on TSA. Triplicate samples were taken at all time points, and the assay was repeated two times.

Survival of recombinant *B. suis* strains in mice. Six-week-old female BALB/c mice (Charles River Laboratories, Wilmington, Mass.) were allowed 1 week of acclimatization. Groups of 7 or 8 mice each were intraperitoneally injected with either 1-2×10<sup>5</sup> CFU of *B. suis* strains 1330, 1330ΔctpA, or VTRS1. Mice were sacrificed at 6 weeks after inoculation and the *Brucella* CFU per spleen determined as described previously (Schurig et al., 1991). Briefly, spleens were collected and homogenized in TSB. Serial dilutions of each spleen's homogenates were plated on TSA plates. The number of CFU that appeared on plates was determined after four days of incubation.

In order to determine the clearance of strains in different time intervals, groups of 25 mice were injected with 0.9- $1.1 \times 10^4$  CFU of *B. suis* strains 1330 or 1330 $\Delta$ ctpA. Groups of five mice injected with each strain were sacrificed at 1, 3,

5, 7 and 9 weeks after inoculation and the *Brucella* CFU per spleen determined as described above.

ELISA. B. suis wild type strain 1330 cells were harvested and killed by boiling for 30 minutes, resuspended at 1:20 in carbonate buffer (pH 9.6) and used to coat the wells of polystyrene plates (100 μl/well; Nunc-Immuno plate with a MaxiSorp surface). After overnight incubation at 4° C., the plates were washed three times in wash buffer (Tris-buffered saline [TBS] at pH 7.4, 0.05% Tween 20) and the diluted mouse serum samples (1:100 dilution in PBS) were added to the wells (100 µl/well). Each serum sample was tested in duplicate wells. The plates were incubated for 2 h at room temperature and washed three times, and isotype-specific goat anti-mouse reagents (Sigma, St. Louis, Mo.) diluted at 1:1,000 in PBS were added to the wells (100 µl/well). After 30 min of incubation at room temperature, the plates were washed three times, and rabbit anti-goat IgG (Sigma) diluted at 1:5,000 in washing buffer (100 µl/well) was added. After 15 min of incubation at room temperature, the plates were washed three times, and 100 µl of substrate solution (TMB Microwell peroxidase substrate; Kirkegaard & Perry Labo- 20 ratories, Gaithersburg, Md.) was added to each well. After 10 min of incubation at room temperature, the enzyme

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mice inoculated with strain  $1330\Delta$ ctpA were intraperitoneally challenged with  $3.2 \times 10^3$  CFU of wild type, virulent *B. abortus* strain 2308. The other five mice injected with PBS and the six mice inoculated with strain  $1330\Delta$ ctpA were challenged with  $1.4 \times 10^5$  CFU of wild type, virulent *B. melitensis* strain 16M. Two weeks post-challenge, mice were sacrificed and the *Brucella* CFU per spleen determined as described above.

Data analyses. Results from the clearance study, ELISA and protection study were analyzed using the Microsoft Excel 2001 program (Microsoft Corporation).

#### Results

Nucleotide sequence of ctpA. At the amino acid level, ctpA gene shared 99% identity with the tail-specific protease of *B. melitensis*. Additionally, it showed up to 77% identity with carboxyl-terminal proteases of a number of bacterial species including *Bartonella Quintana*, *Mesorhizobium loti*, *Sinorhizobium meliloti* and, *Agrobacterium tumefaciens*, and up to 61% identity with a periplasmic protease of other bacteria including *Rhodopseudomonas palustris*, *Rhodobacter sphaeroides*, and *Magnetospirillum magnetotacticum* (Table 2).

TABLE 2

Amino acid level identity of B. suis Cxtp protein to the protein sequences in GenBank				
Bacterial species	Protein	Identity to B. suis CtpA (%)	GenBank Accession #	
B. melitensis	Tail-specific proteinase	99	NP_539132.1	
Mesorhizobium loti	Carboxyl-terminal protease	77	NP_104979.1	
Bartonella quintana	Carboxyl-terminal protease	72	Q44879	
Agrobacterium tumefaciens	Carboxyl-terminal protease	71	NP_355704.1	
Sinorhizobium meliloti	Carboxyl-terminal protease	71	NP_387272.1	
Bradyrhizobium japonicum	Carboxyl-terminal protease	59	NP_771462.1	
Pseudomonas species	Carboxyl-terminal protease	52	NP_747159.1	
Escherichai coli	Carboxyl-terminal protease	31	D00674.1	
Rhodopseudomonas palustris	Periplasmic protease	61	ZP_00009772.1	
Rhodobacter sphaeroides	Periplasmic protease	61	ZP_00007601.1	
Magnetospirillum magnetotacticum	Periplasmic protease	53	ZP_00054906.1	
Azotobacter vinelandii	Periplasmic protease	50	ZP_00089764.1	
Microbulbifer degradans	Periplasmic protease	50	ZP_00065626.1	

4:

reaction was stopped by adding  $100 \mu l$  of stop solution (0.185M sulfuric acid), and the  $A_{450}$  was recorded with a microplate reader (Molecular Devices, Sunnyvale, Calif.).

Protective efficacy of *B. suis* mutant 1330ΔctpA. Sixweek-old female BALB/c mice (The Jackson Laboratory, Bar Harbor, Me.) were allowed 1 week of acclimatization. Groups of 7 mice each were intraperitoneally injected with PBS, strain 1330ΔctpA or strain VTRS1. Two doses of strains 1330ΔctpA or VTRS1 were used in vaccination, i.e., a high dose, which was similar to the dose used in clearance study and the low dose, which was 1 log<sub>10</sub> CFU lower than the above dose. Eight weeks post-inoculation, mice were intraperitoneally challenged with 4.8×10<sup>4</sup> CFU of wild type, virulent *B. suis* strain 1330. Two weeks post-challenge, mice were sacrificed and the *Brucella* CFU per spleen determined as described above.

In a separate trial, six-week-old female BALB/c mice (The Jackson Laboratory, Bar Harbor, Me.) were allowed 1 week of acclimatization. Ten of these mice were intraperitoneally injected with PBS, and another 12 mice were inoculated with 2.1×10<sup>5</sup> CFU of strain 1330ΔctpA. Six weeks post-inoculation, five mice injected with PBS and six

Genomic characterization of CtpA-deficient *B. suis* strain. The primers used to amplify the ctpA gene yielded a 1408-bp product from wild type *B. suis* strain 1330, and a 2189-bp product from strain 1330ΔctpA (FIG. 4). These results indicated that due to double-crossover event, a 471-bp region was deleted from ctpA gene, and the 1252-bp Kan was inserted at the deletion site of strain 1330 genome. A PCR assay with this primer pair produced a 1408-bp band not only from wild type *B. suis*, but also from *B. abortus*, *B. canis* and *B. melitensis* (data not shown).

Growth rates of recombinant *B. suis* strains. The Prcdeficient *E. coli* strain exhibited salt-sensitive and thermosensitive growth (Hara et al., 1991). We investigate if the CtpA-deficient *B. suis* strain exhibits similar growth patterns. After three days of growth on regular TSA plates, colonies of strain 1330ΔctpA appeared approximately one third to half the size of the colonies of strain 1330 (data not shown). In regular LB broth, strain 1330ΔctpA grew approximately 50% slower than strain 1330 did (FIG. 5A) and exhibited no growth when incubated in salt-free LB broth (FIG. 5B). The growth rates of wild type versus mutant strains did not differ as a function of temperature, i.e., wild

type and mutant strains grew at approximately similar rates at 30, 37 or 42° C. (data not shown). Colonies of strain 1330 $\Delta$ ctpA complemented with ctpA appeared equal in size to those of strain 1330 on TSA plates (data not shown). The growth rate of the complemented  $\Delta$ ctpA strain in regular or salt-free LB media was similar to the wild type strain 1330 (FIGS. 5A and 5B).

Complementation of CtpA and Prc activity in CtpA- or Prc-deficient strains. The absence of growth in salt-free media of strain 1330 $\Delta$ ctpA was reverted when the ctpA gene was introduced into this strain (resulting strain 1330 $\Delta$ ctpA [pBB $\Delta$ ctpA]). However, the salt sensitive growth of Prc-deficient *E. coli* strain JE7929 could not be reverted when *B. suis* ctpA gene was introduced into this strain.

Any leakage of periplasmic proteins. In Prc-deficient *E. coli* strain, significant amounts of RNase I and periplasmic alkaline phosphatase were leaked into the culture media (Hara et al. 1991). In order to find out if a similar phenomenon takes place in the CtpA-deficient *B. suis*, we precipitated the protein culture supernatant with acid, and used in SDS/PAGE and Western assays. In either assay, no significant differences were observed between the wild type strain 1330 and the CtpA-deficient strain 1330ΔctpA. No visible protein bands were seen on Western immunoblots with hyper immune anti-*Brucella* goat serum (data not shown), indicating that disruption in ctpA gene may not cause proteins to leak out of cells at significant level.

Phenotypic characterization of recombinant strains. We studied if the mutations in the ctpA gene made any effect on the proteins involved in lipopolysaccharide transport by assessing possible alterations in smooth phenotype. Similar to strain 1330, strain 1330ΔctpA did not retain crystal violet stain, indicating that both these strains possess a smooth phenotype (Table 3). In contrast, rough colonies of strain VTRS1 (Winter et al, 1996) retained crystal violet stain, confirming that these staining results were being correctly interpreted.

Cell morphology. Hara et al., (1991) reported that the Prc-deficient *E. coli* strain acquired a filamentous cell morphology when this strain was introduced into the salt-deficient growth media. We examined if a similar phenomenon takes place in the CtpA-deficient *B. suis*. Gramstaining results revealed that the CtpA-deficient strain did not produce a filamentous phenotype when it was grown in salt-deficient media (data not shown). However, we did not attempt to observe if any other morphological changes 45 occurred, i.e., size of cells, or deformation of cells.

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Persistence of *B. suis* strains in J774 macrophages. To study the attenuation characteristics of ΔctpA *B. suis* strain, the persistence of this strain in J774 mouse macrophage cells was studied (Table 3). At 24 and 48 hours post-inoculation, respectively 5.37 and 5.29 log<sub>10</sub> CFU of live *Brucella* were recovered from strain 1330, and 2.28 and 5.01 log<sub>10</sub> CFU were recovered from strain 1330ΔctpA. This reflects 3.09 and 0.28 log<sub>10</sub> decline of persistence of strain 1330ΔctpA compared with strain 1330, indicating that mutation in ctpA gene makes *B. suis* less persistent in J774 macrophages.

Survival in mice of the *B. suis* strains. To study the attenuation characteristics of ΔctpA *B. suis* strain, BALB/c mice were intraperitoneally inoculated with 5.0-5.3 log<sub>10</sub> CFU, and spleen CFU were determined 6 weeks postinoculation (Table 4). The virulent wild type strain 1330 persisted in mice for more than 6 weeks with only 0.83 log<sub>10</sub> CFU decline, whereas, strain 1330ΔctpA declined by 3.21 log<sub>10</sub> CFU during the same period. In comparison, splenic recovery of attenuated, rough *B. suis* strain VTRS1 declined 2.92 log<sub>10</sub> CFU.

TABLE 4

Clearance from mouse spleens of *B. suis* strains. Six-weeks old BALB/c mice were intraperitoneally inoculated with 5.0-5.3 log<sub>10</sub> CFU, and spleen CFU were determined 6 weeks postinoculation.

ı	Strain	Injected dosage (log <sub>10</sub> CFU/mouse)	CFU 6 weeks after inoculation (Mean + SE $\log_{10}$ /spleen)	Spleen size*
	1330 (wild)	5.24	$4.41 \pm 0.18$	Enlarged
	1330∆ctpA	5.25	$2.04 \pm 0.89^{a}$	Normal
	VTRS1	4.97	$2.05 \pm 1.08^{b}$	Normal

\*Spleen size of mice that were not infected with any bacteria was considered normal.

<sup>a</sup>Completely cleared in one out of eight mice <sup>b</sup>Completely cleared in one out of seven mice

In a separate trial, the splenic clearance of strains was estimated in every two-week intervals. In this work, mice were intraperitoneally inoculated with 4.0-4.1 log<sub>10</sub> CFU of strains 1330 or 1330ΔctpA, and spleen CFU were determined 1, 3, 5, 7, and 9 weeks post-inoculation (FIG. 6). One week after inoculation, the average splenic recovery of the

### TABLE 3

B. suis strains - genes interrupted by mutation, phenotype, and clearance from mouse macrophage 1774 cell lines. 1774 cells were inoculated with B. suis strains. The recovery of strains 24 and 48 hours post-inoculation was determined.

Recovery of Brucellae
from macrophages
(log., CFU/well)

Strain	Gene interrupted by knockout mutagenesis	Phenotype <sup>a</sup>	24-hours of incubation	48-hours of incubation
1330 (wild) 1330∆ctpA	— carboxyl-terminal	Smooth Smooth	$5.37 \pm 0.78$ $2.28 \pm 0.21$	5.29 ± 0.34 5.01 ± 0.15
VTRS1	protease (ctpA) mannoseyltransferase (wboA)	Rough	_	_

<sup>&</sup>lt;sup>a</sup>Assessed with crystal violet colony staining

strain 1330ΔctpA remained 4.0 log<sub>10</sub> CFU, while it was 2.1 log<sub>10</sub> CFU higher in the strain 1330. The persistence of the strain 1330∆ctpA declined faster than that of the strain 1330.

Nine weeks post-inoculation, the CtpA-deficient strain cleared from spleens but strain 1330 was still present. It is noted that the SD values for the CptA mutant at 5 and 7 week post-inoculation (FIG. 6) were quite large because this strain had been completely cleared from the spleen of at least one mouse (i.e. 0 CFU).

Induction of immune responses in mice. Specific antibody responses of the vaccinated mice were determined by ELISA (FIG. 7). Mice injected with saline produced negligible amounts of IgG1 or IgG2a recognizing B. suis antigens. Compared to the sera from mice vaccinated with B. abortus 15 strain RB51 (obtained from A. Contreras, Virginia Tech), sera from mice vaccinated with strain 1330ΔctpA contained about seven fold greater IgG1 (P<0.001). Nevertheless, sera from 1330∆ctpA-vaccinated mice contained only slightly higher IgG2a levels than sera from RB51-vaccinated mice 20 (P<0.01).

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#### TABLE 5

Protection induced by recombinant B. suis strains against challenge with B. suis virulent strain 1330. Six weeks old BALB/c mice were injected with PBS or B. suis strains as shown. Six weeks post-injection, the mice were challenged with 4.7 Log<sub>10</sub> CFU of B. suis strain 1330. Spleen CFU were determined two weeks post-challenge

)	Vaccine strain	Dose injected log <sub>10</sub> CFU	Recovery of strain 1330 from spleens log <sub>10</sub> CFU	Units of protection log <sub>10</sub> CFU	Spleen size*
	PBS	_	5.90 ± 0.24	_	Larger than normal
	VTRSI	Low dose 4.20	$5.91 \pm 0.54$	-0.01	Larger than normal
		High dose 5.20	$4.64 \pm 0.39$	1.26	Normal
	1330∆ctpA	Low dose 4.34	$2.70 \pm 0.55$	3.20	Normal
		High dose 5.34	$2.15 \pm 0.96$	3.75	Normal

\*Spleen size of mice that were not infected with any bacteria was considered normal.

TABLE 6

Protection induced by recombinant B. suis strain 1330ΔctpA against challenge with B. abortus virulent strain 2308 and B. melitensis virulent strain 16M. Six weeks old BALB/c mice were injected with PBS or B. suis strain 1330ΔctpA as shown. Six weeks post-injection, the mice were challenged with B. abortus strain 2308 or B. melitensis strain 16M, and spleen CFU were determined two weeks post-challenge

Inoculation	Dose injected log <sub>10</sub> CFU/mouse	Challenge strain	Challenge dose log <sub>10</sub> CFU/mouse	Recovery of challenge strains from spleens log <sub>10</sub> CFU/mouse	Units of protection log <sub>10</sub> CFU/mouse
PBS	_	2308	3.51	5.03 ± 0.07	4.71
1330∆ctpA	5.332	2308	3.51	$0.32 \pm 0.78^{a}$	
PBS	_	16M	5.14	$5.62 \pm 0.29$	0.37
1330∆ctpA	5.32	16M	5.14	$5.25 \pm 0.31$	

aCompletely cleared in five out of six mice

Protective efficacy of attenuated B. suis strains. Mice 45 Discussion immunized with 4.34 and 5.34 log<sub>10</sub> CFU of strain 1330∆ctpA demonstrated 3.20 and 3.75 log<sub>10</sub> units of protection, respectively (Table 5). All colonies harvested from spleens of mice injected with this strain were sensitive to 50 kanamycin, indicating that they all were from the challenge strain 1330 (kanS), as opposed to the vaccine strain 1330ΔctpA (kanR). In comparison, strain VTRS1 provided no protection when mice were vaccinated with 4.20 log<sub>10</sub> CFU, but provided 1.26 log<sub>10</sub> CFU protection when vaccinated with 5.20 log<sub>10</sub> CFU dose. Nearly one-quarter of the colonies harvested from mice immunized with  $5.20 \log_{10}$ CFU of strain VTRS1 were resistant to kanamycin indicating that the VTRS1 dose had not been completely cleared from the spleens in 10 weeks. In a separate trial, it was shown that immunization with strain 1330ΔctpA induced 4.71 and 0.37 log<sub>10</sub> CFU units of protection against challenge with strains 2308 and 16M, respectively (Table 6).

The deduced amino acid sequence from ctpA gene showed substantial homology with the Ctps of a number of bacterial species. In addition, this deduced amino acid sequence showed considerable homology with the periplasmic proteases of other related bacterial species. The B. suis CtpA showed 31% homology at the amino acid level to Prc protein identified as carboxyl-terminal processing protease for PBP 3 of E. coli (Hara et al., 1991; Silber et al., 1992). Cell fractionation studies had indicated that Prc is localized in the periplasmic space of E. coli (Hara et al., 1991). Based on the greater homology between bacterial Ctps and periplasmic proteases we believe that these two protein groups are the same, even though they had been named differently. It is possible that B. suis CtpA is involved in carboxylterminal processing protease activities and is located in the periplasmic space of the cell. This possibility awaits further experimental confirmation.

Strains 1330\DeltactpA and 1330 possessed a smooth phenotype. Apparently, the mutation of ctpA gene does not have-

impact on the transport of O-side chain to the outer membrane. Again this observation is consistent with CtpA affecting the processing of proteins as opposed to carbohydrates. Strain 1330\(\Delta\)ctpA produced relatively smaller colonies on TSA plates, and exhibited slower growth in regular growth media suggesting that the function of CtpA is important for the growth and cell division of B. suis. Zero growth of strain 1330ΔctpA in salt-free media suggests that ctpA function involves either directly or indirectly protection of the cell from osmotic stresses. Hara et al., (1991) observed that Prc-deficient E. coli did not grow in salt-free media. Apparently, in E. coli and in B. suis respectively, Prc and CtpA functions are dispensable for growth in a normal osmolar milieu. The reduced growth in salt-free media exhibited by CtpA-deficient mutant mimics that seen for the Prc-deficient mutant of E. coli (Hara et al., 1991) and further suggests that the B. suis CtpA is a protease. Prc-deficient E. coli grew in salt-free media at low temperatures (30° C.) but not at high temperatures (42° C.) exhibiting temperature dependency. In contrast, CtpA-deficient B. suis strain did not grow at any temperature when it was introduced into salt-free media, 20 indicating that CtpA in Brucella does not have a temperature dependency.

Hara et al. (1991) reported that when the Prc-deficient *E. coli* cells were introduced into the salt-free media, those cells exhibited a filamentous phenotype. These workers attributed this phenomenon to the interruption of the processing/maturation of PBP 3 protein in Prc-deficient strain. The present work shows that the cells of the CtpA-deficient *B. suis* did not exhibit a filamentous shape when they were introduced into the salt-free growth media. This result suggests that CtpA may not be involved in the processing of PBP 3 of *B. suis*. However, this possibility awaits further experimental confirmation.

Colonies produced by strain 1330ΔctpA complemented with the ctpA gene were similar in size to those produced by wild type strain 1330. Additionally, the growth pattern of this strain was similar to that of strain 1330. These results indicate that complementation of ctpA gene restored the CtpA activity of mutant 1330ΔctpA. These results further suggest that the phenotype of strain 1330ΔctpA is the result of a specific mutation in ctpA and not a polar effect.

Hara et al., (1991) reported that disruption of prc gene of *E. coli* resulted in leakage of proteins from cells. However, the mutation in ctpA gene of *B. suis* did not cause any leakage of proteins, suggesting that this mutation may not affect the integrity of the cell wall.

When the ctpA gene was introduced into Prc-deficient *E. coli*, its growth in salt-free media could not be restored. This may be due to significant structural differences between the CtpA and Prc proteins. These proteins shared only 31% homology at the amino acid level.

Overall, the Prc and CtpA protein appear to regulate the salt-sensitive growth of *E. coli* and *B. suis*. However, unlike that of Prc, the activity of CtpA is not dispensable at different growth temperatures. Additionally, it is apparent that, contrary to Prc, CtpA is not involved in processing of the PBP that regulates murein synthesis/cell division, or retention of periplasmic proteins like RNaseI or alkaline phosphatase.

When grown in J774 macrophages, the persistence of strain 1330ΔctpA declined significantly after 24 hours of incubation, indicating that CptA is important for survival of *B. suis*, particular against early killing by macrophages and neutrophils. The clearance studies in mice revealed that one week after inoculation, significantly lower numbers of strain 1330ΔctpA were recovered than strain 1330. Nine weeks after inoculation, strain 1330ΔctpA was cleared completely from mouse spleens whereas strain 1330 was still present. These findings indicate that mutation in the ctpA gene makes *B. suis* attenuated. Overall, the slow growth of this mutant

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strain in enriched media, and its low persistence in mice and mouse macrophages suggest that it has a diminished capacity for extracellular and intracellular growth.

Brucella species are intracellular pathogens, and therefore, cell-mediated immune (CMI) responses are critically important in preventing brucellosis (Pavlov et al., 1982; Zhan et al., 1993; Murphy et al., 2001). Cattle brucellosis vaccine B. abortus strain RB51 induces preferentially Th1 type associated CMI responses and protects cattle against Brucella infections (Vemulapalli et al., 2000a; Vemulapalli et al., 2000 b). In vaccinated mice, strain 1330ΔctpA produced IgG2a at a slightly higher level than did strain RB51. In addition, unlike strain RB51, strain 1330ΔctpA also induced a greater level of IgG1. Overall, the IgG1 and IgG2a antibody responses induced by strain 1330ΔctpA appear more balanced than those induced by strain RB51.

In the protection study, all colonies harvested from mice vaccinated with strain 1330ΔctpA were sensitive to kanamycin indicating that this strain has been completely cleared from mouse spleens during the 10-week immunization period. Strain 1330\(Delta\text{train}\) induced excellent protection against challenge with B. suis strain 1330, and the level of protection increased slightly with an increased vaccine dose. The protection induced by strain 1330∆ctpA was much greater than that induced by rough B. suis strain VTRS1. The superior protection against 1330 challenge by the strain 1330∆ctpA compared with that elicited by strain VTRS1 can be attributed to two reasons: (A) Strain 1330ΔctpA being a smooth strain may be inducing antibody responses to O-side chain. This is likely to reflect published literature suggesting that the specific humoral and cellular responses to the O-side chain are important in producing good protection (Araya et al., 1989; Araya and Winter, 1990; Corbel, 1997). (B) Strains 1330 and 1330∆ctpA belong to biovar-1 whereas, VTRS1 belongs to biovar-4. Apparently, a biovar-4 vaccine is not as efficient as a biovar-1 vaccine against challenge with biovar-1 challenge. Within a single species of Brucella (e.g. Brucella suis) different groups exhibiting expression of different antigens (protein types of sugar types) are described as "biovars". For instance, Brucella suis biovar-1 and biovar-4 cause brucellosis in swine in other animals. The bacteria in these two groups share many similarities, but they differ to some degree in terms of the expression of specific antigens.

Strain 1330ΔctpA also induced excellent protection against challenge with *B. abortus* strain 2308, indicating that this strain is effective in providing protection against challenge with heterologous *Brucella* species. Nevertheless, strain 1330ΔctpA did not induce satisfactory protection against challenge with *B. melitensis* strain 16M. The less protective efficacy of strain 1330ΔctpA against 16M challenge may be attributable to the structural differences of O-side chain between *B. melitensis* and *B. suis*. Recent work has revealed that monoclonal antibodies to *B. abortus* O-side chain (Bru-38) do not recognize the O-side chain of *B. melitensis*, suggesting that the O-side chain of other *Brucella* species may not induce protection against *B. melitensis* challenge (unpublished data).

As a vaccine candidate, strain 1330ΔctpA possesses a number of advantages over the other brucellosis vaccines. The currently used vaccine candidates, *B. abortus* strains RB51 (Schurig et al., 1991), S19 (Nicoletti, 1990), and *B. melitensis* strain Rev1 (Alton et al., 1967), were developed through laboratory selection procedures, and therefore, their genetic makeups are not known. Even though they induce substantial protection against brucellosis in animals, vaccination with strains S19 (Manthei and Beckett) or Rev1 (Blasco and Bardenstein) sometimes result in abortion. Strains S19 (Young 1983) and Rev1 (Blasco, and Diaz, 1993; Young 1983) are pathogenic to humans. Strain RB51

is resistant to the antibiotic rifampicin, one of the very few antibiotics available for treatment of brucellosis in humans (Joint FAO/WHO, 1986). In contrast, strain 1330ΔctpA was developed by knockout mutagenesis and its genetic make up is well defined. Further, it induces excellent protection 5 against challenge with B. abortus strain 2308 or B. suis strain 1330.

#### Example 2

#### Electron Microscopy Studies

When observed with the electron microscope, wild type Brucella suis (strain 1330) cells possessed their native coccobaccillus cell morphology. No difference in cell morphology was seen between strain 1330 cells grown in growth media with salt (FIG. 8) or without salt (FIG. 9). Additionally, these cells possessed the typical ultrastructure of Brucella cells, namely, the outer membrane, periplasmic space, and cytoplasmic membrane.

However, the invented strain 1330 ctpA exhibited a spherical cell morphology when grown in media with salt. The cell diameter also appeared to be increased slightly. The outer membrane was partially separated from some of the cells (FIG. 10). When grown in media without salt, the membrane dissociated from the rest of the cell, and the cell morphology was significantly altered (FIG. 11).

In other bacteria (i.e., Escherichia coli and Bacillus subtilis), when the expression or processing of Penicillin-Binding Proteins-1 and -2 (PBP-1 and PBP-2) is inhibited, the cells are known to acquire a spherical cell morphology, and the cell diameter increases. Accordingly, it is likely that in the invented strain 1330 ctpA, the expression or processing of PBP-1 and/or PBP-2 has been altered due to mutation of the CtpA protein.

Further, in other bacteria, PBP enzymes are involved in 35 synthesis of the cell wall peptidoglycan layer. Therefore, the dissociation of the cell membrane from the rest of the cell and the loss of cell integrity of strain 1330 ctpA is likely attributable to alteration of the functions of PBP-1 and PBP-2 enzymes due to mutation of the ctpA gene.

#### REFERENCES FOR EXAMPLES 1 AND 2

- Acha, P., and B. Szyfres. 1980. Zoonoses and communicable diseases common to man and animals, p.28-45. In Pan 45 Keiler, K. C., P. R. H. Waller, and R. T. Sauer. 1996. Role of American Health Organization, Washington, D.C.
- Alton, G. G., L. M. Jones, and D. E. Pietz. 1975. Laboratory techniques in Brucellosis. World Health Organization monograph series no. 55. In World Health Organization, Geneva, Switzerland.
- Altschul, S., W. Gish, W. Miller, E. Myers and D. Lipman. 1990. Basic local alignment search tool. J. Mol. Biol. 215:403-410.
- Anderson, T. D., N. F. Cheville, and V. P. Meador. 1986. 55 McQuiston, J. R., G. G. Schurig, N. Sriranganathan, and S. Pathogenesis of placentitis in the goat inoculated with Brucella abortus. II. Ultrastructural studies. Vet. Pathol. 23:227-239.
- Araya, L. N., and A. J. Winter. 1990. Comparative protection of mice against virulent and attenuated strains of Brucella abortus by passive transfer of immune T cells or serum. Infect. Immun. 58:254-256.
- Araya, L. N., P. H. Elzer, G. E. Rowe, F. M. Enright, and A. J. Winter. 1989. Temporal development of protective cell-mediated and humoral immunity in BALB/c mice infected with Brucella abortus. J. Immunol. 143:3330-3337.

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- Arenas, G. N., A. S. Staskevich, A. Aballay, and L. S. Mayorga. 2000. Intracellular trafficking of Brucella abortus in J774 macrophages. Infect. Immun. 68:4255-4263.
- Baldwin, C. L., and A. J. Winter. 1994. Macrophages and Brucella. Immunol. Ser. 60:363-380.
- Chung, S., and D. A. Bryant. 1992. p 69-72. In N. Murata (ed.), Research in Photosynthesis, vol. I.
- Corbel, M. J. 1997. Brucellosis: an overview. Emerg. Infect. Dis. 3:213-221.
- 10 Corbel, M. J., and W. J. Brinley Morgan. 1984. Genus Brucella Meyer and Shaw 1920, 173 AL. p. 377-390. In N. R. Krieg and J. G. Holt (ed.), Bergey's Manual of Systematic Bacteriology, vol. 1. Williams and Wilkins, Baltimore.
- 15 Dalbey, R. E., and G. von Heijne. 1992. Signal peptidases in prokaryotes and eukaryotes—a new protease family. Trends Biochem. Sci. 17:474-478.
  - Francis M. S., and C. J. Thomas. 1997. f1 Mutants in the CtpA copper transporting P-type ATPase reduce virulence of Listeria monocytogenes. Microbial Pathogenesis 22:67-78.
  - Hara, H., Y. Yamamoto, A. Higashitani, H. Suzuki, and Y. Nishimura. 1991. Cloning, mapping, and characterization of the Escherichia coli prc gene, which is involved in C-terminal processing of penicillin-binding protein 3. J. Bacteriol. 173:4799-4813.
  - Inagaki, N., Y. Yamamoto, H. Mori, and K. Satoh. 1996. Carboxyl-terminal processing protease for the D1 precursor protein: cloning and sequencing of the spinach cDNA. Plant Mol. Biol. 30:39-50.
  - Islam, M. R., J. H Grubb, and W. S. Sly. 1993. C-terminal processing of human beta-glucuronidase. The propeptide is required for full expression of catalytic activity, intracellular retention, and proper phosphorylation J. Biol. Chem. 268:22627-22633
  - Jimenez de Bagues, M. P., C. M. Martin, and J. M. Blasco. 1992. An ELISA with Brucella lipopolysaccharide antigen for the diagnosis of B. melitensis infection in sheep and for the evaluation of serological responses following subcutaneous or conjunctival B. melitensis strain Rev 1 vaccination. Vet Microbiol. 30:233-241.
  - Keiler, K. C., and R. T. Sauer. 1998. p. 460-461 In A. J. Barrett, N. D. Rawlings, and J. F. Woessner (ed.), Handbook of Proteolytic Enzymes, Academic Press, London.
  - a peptide tagging system in degradation of proteins synthesized from damaged messenger RNA. Science 271: 990-993
  - Kovach, M. E., R. W. Phillips, P. H. Elzer, R. M. Roop II, and K. M. Peterson. 1994. pBBR1 MCS: a broad-host range cloning vector. BioTechniques 16:800-802.
  - Laemmli, U. K. 1970. Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature 227:680-685.
- M. Boyle. 1995. Transformation of Brucella species with suicide and broad host-range plasmids. Methods Mol. Biol. 47:143-148.
- Menon, N. K., J. Robbins, M. D. Vartanian, D. Patil, H. D. Peck, A. L. Menon, R. L. Robson, and A. E. Przybyla. 1993. Carboxy-terminal processing of the large subunit of [NiFe] hydrogenases. FEBS Lett. 331:91-95.
- Murphy, E. A., J. Sathiyaseelan, M. A. Parent, B. Zou and C. L. Baldwin. 2001. Interferon-gamma is crucial for surviving a Brucella abortus infection in both resistant C57 BL/6 and susceptible BABB/c mice. Immunology 103: 511-518.

- Naroeni, A., N. Jouy, S. Ouahrani-Bettache, J. P. Liautard, and F. Porte. 2001. Brucella suis-impaired specific recognition of phagosomes by lysosomes due to phagosomal membrane modifications. Infect. Immun. 69:486-493.
- National Academy Press. 1977. Brucellosis research: an 5 evaluation. p. 61-77. In Report of the subcommittee on Brucellosis research. National Academy Press, Washington, D.C.
- Oelmüller, R., R. G. Herrmann, and H. B. Pakrasi. 1996. Molecular studies of CtpA, the carboxyl-terminal pro- 10 cessing protease for the D1 protein of the photosystem II reaction center in higher plants. J. Biol. Chem. 271: 21848-21856.
- Paetzel, M. and R. E. Dalbey. 1997. Catalytic hydroxyl/ amine dyads within serine proteases. Trends Biochem. Sci. 22:28-31.
- Pakrasi, H. B. 1998. p. 462-463. In A. J. Barrett, N. D. Rawlings, and J. F. Woessner (ed.), Handbook of Proteolytic Enzymes, Academic Press, London.
- Paulsen, I. T., R. Seshadri, K. E. Nelson, J. A. Eisen and S. M. Boyle. 2002. The Brucella suis genome reveals fundamental similarities between animal and plant pathogens and symbionts. PNAS 99:13148-13153.
- Pavlov, H., M. Hogarth, I. F. C. McKenzie and C. Cheers. 25 1982. In vivo and in vitro effects of monoclonal antibody to Ly antigen on immunity to infection. Cell. Immunol. 71:127-138.
- Pizarro-Cerda, J., S. Meresse, R. G. Parton, G. van der Goot, Gorvel. 1998. Brucella abortus transits through the autophagic pathway and replicates in the endoplasmic reticulum of nonprofessional phagocytes. Infect. Immun. 66:5711-5724.
- Ried, J. L., and A. Colmer. 1987. An npt-sacB-sacR car- 35 tridge for constructing directed, unmarked mutations in gram-negative bacteria by marker exchange-eviction mutagenesis. Gene. 57:239-246.
- Sambrook, J., E. F. Fritsch and T. Maniatis. 1989. Molecular cloning: a laboratory manual, 2 nd ed. Cold Spring Harbor 40 Laboratory Press, Cold Spring Harbor, N.Y.
- Satoh, K. 1998, p. 463-464. In A. J. Barrett, N. D. Rawlings, and J. F. Woessner (ed.), Handbook of Proteolytic Enzymes, Academic Press, London.
- Schurig, G. G., R. M. Roop, T. Bagchi, S. M. Boyle, D. 45 Buhrman, and N. Sriranganathan. 1991. Biological properties of RB51: a stable rough strain of Brucella abortus. Vet. Microbiol. 28:171-188.
- Silber, K. R., K. C. Keiler, and R. T. Sauer. 1992. Tsp: a 50 tail-specific protease that selectively degrades proteins with nonpolar C termini. Proc. Natl. Acad. Sci. U. S. A.
- Theroux, S. J., T. E. Redlinger, R. C. Fuller, and S. J. Robinson. 1990. Gene encoding the 5.7-kilodalton chlo-55 rosome protein of Chloroflexus aurantiacus: regulated message levels and a predicted carboxy-terminal protein extension. J. Bacteriol. 172:4497-504.
- Vemulapalli, R., A. J. Duncan, S. M. Boyle, N. Sriranganathan, T. E. Toth, and G. G. Schurig. 1998. Cloning and 60 sequencing of yajC and secD homologs of Brucella abortus and demonstration of immune responses to YajC in mice vaccinated with B. abortus RB51. Infect. Immun. 66:5684-5691.
- Vemulapalli R., Y. He, L. S. Buccolo, S. M. Boyle, N. 65 Sriranganathan, G. G. Schurig. 2000a. Complementation of Brucella abortus RB51 with a functional wboA gene

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- results in O-antigen synthesis and enhanced vaccine efficacy but no change in rough phenotype and attenuation. Infect. Immun. 68:3927-3932.
- Vemulapalli R., Y. He, S. Cravero, S. M. Boyle, N. Sriranganathan, and G. G. Schurig. 2000b. Overexpression of protective antigen as a novel approach to enhance vaccine efficacy of Brucella abortus strain RB51. Infect. Immun. 68:3286-3289.
- White, P. G., and J. B. Wilson. 1951. Differentiation of smooth and nonsmooth colonies of Brucellae. J. Bacteriol. 61:239-240.
- Winter, A. J., G. G. Schurig, S. M. Boyle, N. Sriranganathan, J. S. Bevins, F. M. Enright, P. H. Elzer, and J. D. Kope. 1996. Protection of BALB/c mice against homologous and heterologous species of Brucella by rough strain vaccines derived from Brucella melitensis and Brucella suis biovar 4. Am. J. Vet. Res. 57:677-683
- Yamamoto, Y., N. Inagaki, and K. Satoh. 2001. Overexpression and characterization of carboxyl-terminal processing protease for precursor D1 protein: regulation of enzymesubstrate interaction by molecular environments. J. Biol. Chem. 276:7518-7525
- Zhan, Y., J. Chang and C. Cheers. 1993. Cytokine response of, T-cell subsets from Brucella abortus infected mice to soluble Brucella proteins. Infect. Immun. 61:2841-2847.

#### Example 3

The carboxyl-terminal protease (CtpA) proteins are a A. Sola-Landa, I. Lopez-Goni, E. Moreno, and J. P. 30 novel family of enzymes. The putative tail-specific protease (TspA) of Brucella melitensis strain 16M is a member of the CtpA family. Sequence analyses predicted that the mature TspA protein is localized in the periplasmic space. Out of total 443 amino acids of the B. melitensis TspA sequence, 174 amino acids are conserved among the CtpA proteins of at least six bacterial species. The Asp238, Arg240, Ser300, Glu303, and Lys325 amino acid residues of B. melitensis TspA sequence correspond with those amino acid residues critical for the catalytic activity of the CtpA protein of Synechocystis species. Among these, Ser300/Lys325 appears to be the catalytic dyad of this protein. A tspA mutant B. melitensis strain (16MΔtspA), generated by allelic exchange, produced smaller colonies on enriched agar plates, and exhibited zero growth in salt-free enriched medium compared to the wild type strain 16M or the complemented mutant strain 16M\Delta\tspA[ctpA<sup>+</sup>]. Western immunoblotting assays revealed that the tspA mutant upregulated the expression of at least two proteins. Electron microscopy revealed that in contrast to the native coccobacillus shape of wild type strain, the tspA mutant possessed a spherical shape with an increased cell diameter. In the J774 mouse macrophage cell line, 24 hours after infection, the survival of the tspA mutant strain declined by approximately  $1.2 \log_{10}$  colony forming units relative to the wild type strain. These observations suggest that the TspA protein is involved in determining growth, protein expression, cell morphology, and intracellular persistence of B. melitensis. Key words: *Brucella melitensis*, carboxyl-terminal protease, Tail-specific protease, morphology, protein expression,

#### INTRODUCTION

intracellular persistence.

Brucellae are gram-negative intracellular bacterial pathogens of both humans and animals (Corbel and Brinley Morgan, 1984). Among the six recognized Brucella species, Brucella melitensis is the main etiologic agent involved in ovine and caprine brucellosis and is also the most pathogenic species for humans (Acha and Szyfres, 1980). The

pathological manifestations of brucellosis are diverse and include arthritis, endocarditis and meningitis in man, whereas animal brucellosis is characterized by spontaneous abortion (Young, 1983). Molecular characterization of cellular factors involved in regulating critical cellular functions would enhance the basic knowledge about physiology of bacteria. Additionally, identifying the factors important for intracellular persistence of *Brucella* will aid development of live attenuated vaccines against this bacterium.

Many proteins destined for extracytoplasmic locations are initially synthesized as precursor forms and processed into mature forms by proteolytic cleavage to remove short peptide sequences, either near the amino terminus or near the carboxyl terminus of such proteins. The endoproteases responsible for cleaving of amino-terminal peptides are called amino-terminal processing proteases and have been identified and studied in a number of systems (Dalbey and Von Heijne, 1992). During recent years, a relatively new class of endoproteases with carboxyl-terminal processing activities has been described in various bacteria and organellar systems. These proteins have been designated as Car- 20 boxyl-Terminal Protease (CtpA) proteins. The CtpA proteins are serine proteases that utilize a Ser/Lys catalytic dyad instead of the well-known Ser/His/Asp catalytic triad (Paetzel and Dalbey, 1997).

As a whole CtpA proteins are not well understood. To 25 date, the best-characterized CtpA is that from the cyanobacterium Synechocystis sp. strain PCC 6803 (Shestakov et al., 1994). This enzyme is responsible for processing of the D1 precursor polypeptide of Photosystem II (Nixon et al., 1992). The CtpA protein is also involved in D1 processing 30 in higher plants (Takahashi et al., 1988), and green algae Scenedesmus obliquus (Trost et al., 1997). Another wellstudied CtpA is the tail-specific protease (Tsp) enzyme from E. coli (Silber et al., 1992). The E. coli Tsp is responsible for cleavage of C-terminal 11 amino acid residues of precursor form of Penicillin-Binding Protein-3 (PBP-3) (Hara et al., 1991). The PBP-3 is believed to involve in determining division of rod-shaped cells in bacteria (Popham and Young, 2003). A mutant E. coli strain deficient in Tsp expression was defective in processing of PBP-3 (Hara et al., 1989, 1991), and exhibited filamentous cell morphology, in con-40 trast to the cocco-bacillus morphology of the wild type E. coli (Hara et al., 1991). Additionally, this mutant showed thermo-sensitive growth on salt-free L-agar plates, suggesting that Tsp was involved in protection of cell from thermal and osmotic stresses (Hara et al., 1991). More recently, Tsp 45 has been shown to recognize and degrade several aberrant proteins with nonpolar C-termini, with strongest preference for alanine, at the three C-terminal residues (Keiler et al.,

A putative CtpA is present in Borrelia burgdorferi strain 50 B31 (Ostberg et al., 2004). Inactivation of the ctpA gene encoding this protein resulted in altered expression pattern of a number of proteins. The integral outermembrane protein P13 and the hypothetical protein BB0323 were identified as the substrates for the CtpA of this bacterium (Ostberg et al., 2004). A tsp homologue that expresses a CtpA protein is present in Salmonella typhimurium. The tsp mutant S. typhimurium had a reduced survival within macrophages suggesting that this gene may play a role in virulence (Baumler et al., 1994). A CtpA protein has also been identified in Bartonella bacilliformis, but no target for this enzyme has 60 been identified yet. The ctpA gene encoding this protease is located immediately upstream of the ialAB locus that confers the bacterium the ability to invade human erythrocytes (Mitchell et al., 1997).

The knowledge about protein processing and protein 65 modification in *Brucella* is limited. We recently reported that *B. suis* produces a putative CtpA protein that is involved in

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regulating the growth, cell morphology and intracellular persistence of this bacterium (Bandara et al., 2005, in press). A homologue of genes encoding CtpA proteins, designated tspA (GenBank locus tag BMEI0214 and Gene ID:1195926) is present on *B. melitensis* 16M chromosome I (Paulsen et al., 2002). In the present communication we report that the tspA gene influences the salt-sensitive growth, protein expression, cell morphology and intracellular persistence of *B. melitensis*.

#### Methods

Bacterial Strains, Plasmids, and Reagents

Brucella melitensis strain 16M was obtained from our culture collection. Brucellae were grown in LB broth (Difco Laboratories, Sparks, Md.) with or without sodium chloride at 37° C. to determine whether growth was osmo-sensitive. For all other assays, brucellae were grown either in trypticase soy broth (TSB) or on trypticase soy agar (TSA) plates (Difco) at 37° C. in the presence of 5% CO<sub>2</sub> as previously described (Schurig et al., 1991). Bacteria containing plasmids were grown in presence of ampicillin or kanamycin at 100 µg/ml concentration as described below. The mouse macrophage-like cell line was J774 obtained from American Type Culture Collection (Manassas, Va.).

All experiments with live brucellae were performed in a Biosafety Level 3 facility at the Infectious Disease Unit of the Virginia-Maryland Regional College of Veterinary Medicine as per standard operating procedures approved by the Centers for Disease Control and prevention.

#### DNA and Protein Sequence Analyses

The nucleotide sequence of tspA gene was analyzed with DNASTAR software (DNASTAR, Inc., Madison, Wis.). The deduced protein sequence of the tspA gene was aligned with the protein sequences of other bacteria using the ClustalV (PAM250) megalign program of the DNASTAR. The sequences of proteins that were more than 59% identical to the TspA were used in this analysis. The SignalP 3.0 Server of the Technical University of Denmark located at the website at cbs.dtu.dk (Bendtsen et al., 2004) was employed to predict the presence of any signal sequence of the putative TspA protein. The destination of the TspA protein upon translation and processing was predicted using the Subloc v1.0 server of the Institute of Bioinformatics of the Tsinghua University at the website located at bioinfo.tsinhua.edu.cn.

Mutation of the B. melitensis tspA Gene by Allelic Exchange

The tspA of B. melitensis shares 99% homology at amino acids level with the ctpA gene encoding putative CtpA protein of B. suis. A suicide vector (pGEMctpAK) that we previously used to mutate the ctpA gene of B. suis (Bandara et al., 2005-in press) was used to mutate the tspA gene of B. melitensis. One microgram of pGEMctpAK was used to electroporate B. melitensis strain 16M with a BTX ECM-600 electroporator (BTX, San Diego, Calif.), as described previously (McQuiston et al., 1995); two colonies of strain 16M were obtained from a TSA plate containing kanamycin (100 µg/ml). These colonies were streaked on a TSA plate containing ampicillin (100 µg/ml) to determine if a single- or double-crossover event had occurred. Both the colonies did not grow on ampicillin containing plates suggesting that a double-crossover event had occurred in them. PCR with a primer pair (forward primer 5 '-GGGGTACCGTGGTG-GACTGA-3 ' (SEQ ID NO: 12) and reverse primer 5 '-GGCTGCAGTCCCGCGTTTTTGTCTT-3 ') (SEQ ID NO: 13) (Ransom Hill Bioscience, Inc., Ramona, Calif.) confirmed that a double-crossover event had taken place in the two transformants. One of these strains was chosen for further analyses and designated 16MΔtspA.

Complementation of tspA Gene Activity in Mutant  $16M\Delta tspA$ 

We had previously cloned the 1.4-kb DNA fragment containing *B. suis* ctpA gene into the broad-host range vector pBBR4MCS (Kovach et al., 1995) to produce recombinant 5 plasmid pBBctpA (Bandara et al., 2005-in press). One microgram of pBBctpA was used to electroporate the tspA mutant *B. melitensis* strain 16MΔtspA; several colonies of strain 16MΔtspA were picked from a TSA plate containing ampicillin (100 μg/ml). One of these colonies was chosen for further analyses and designated as 16MΔtspA[ctpA<sup>+</sup>].

Denaturing Gel Electrophoresis of Proteins and Western Blotting

The wild type B. melitensis strain 16M, the tspA mutant 16MΔtspA and the complemented tspA mutant 16MΔtspA [ctpA<sup>+</sup>] were grown on TSA plates for four days to stationary growth phase. The cells were harvested from plates and resuspended in Laemmli sample buffer (Boston BioProducts Inc., Worcester, Mass.), boiled for 20 min and electrophoresed on 15% SDS/PAGE gels according to standard proce- 20 dures (Laemmli, 1970). Gels containing the separated proteins were either stained with Coomassie brilliant blue G (Sigma Chemical Co.) or used for Western blot analysis. Western blotting was performed as previously described (Vemulapalli et al., 1998). Briefly, proteins separated by SDS-PAGE were transferred to a nitrocellulose membrane by using a Trans-blot semidry system (Bio-Rad Laboratories, Hercules, Calif.). For analysis of total proteins, the membranes were incubated with polyclonal serum obtained from a goat hyperimmunized with B. abortus strain RB51 (Goat-48) (Roop et al., 1992). After 24 hours of incubation with primary sera, the blots were developed with anti-goat IgG (whole molecule) conjugated with horseradish peroxidase (Sigma Chemical Co).

#### Growth Rates of B. melitensis Strains

Salt-free LB media was prepared by mixing bactotryptone and yeast extract in water per manufacturer's instruction (Difco), but omitting sodium chloride. Single colonies of the wild type, the tspA mutant, and the complemented tspA mutant strains were grown at 37° C. for 72 hours to stationary phase in 5 ml of TSB. The cells were harvested by centrifugation, and resuspended in distilled water. The cell suspensions were used to inoculate 15 ml of LB broth or salt-free LB broth in Klett side-arm flask to 6 to 21 Klett units. Cultures were grown at 37° C. at 200 rpm; Klett readings were recorded every two hours in a Klett-Summerson calorimeter (Klett-Summerson, New York, N.Y.).

#### Electron Microscopy

The wild type and the tspA mutant strains were grown in 25 ml LB broth to stationary phase. The cells were harvested 50 by centrifugation at 2000×G for 15 min and introduced to 100 ml LB broth. The cultures were incubated overnight at 37° C. with vigorous shaking. The cells were harvested by centrifugation at 2000×G for 15 min, and fixed overnight at 4° C. in formadehyde-paraformaldehyde in cacodylate buffer (Banai et al., 2002). The samples were then processed for thin section electron microscopy as described by Banai et al., (2002). The sections were mounted on copper grids, stained with uranyl acetate and lead citrate, and examined with a JOEL 100 CX-II transmission electron microscope (Zeiss 10 C, Carl Zeiss Inc., New York, N.Y.) at 10,000× magnification.

Persistence of Recombinant B. melitensis Strains in Macrophages

TSA plates were inoculated with single colonies of B. 65 *melitensis* strains. After four days of incubation at 37° C. with 5% CO<sub>2</sub>, the cells were harvested from plates, washed

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with PBS, resuspended in 20% glycerol, and frozen at -80° C. The number of viable cells was determined after dilutions of the cell suspensions were plated on TS agar. The mouse macrophage-like cell line J774 was seeded at a density of 5×10<sup>5</sup>/ml in Dulbecco's modified essential medium (DMEM) (Fisher Scientific, Atlanta, Ga.) into 24-well tissue culture dishes and cultured at  $37^{\circ}$  C. with 5% CO<sub>2</sub> until confluent. The tissue culture medium was removed, 200 µl  $(2\times10^8 \text{ cells})$  of the bacterial suspension in PBS was added, and the cells were incubated at 37° C. for four hours. The suspension above the cell monolayer was removed, and the cells were washed three times with PBS. One milliliter of DMEM containing 25 µg of gentamicin was added, and the cells were incubated for 24 hours at 37° C. At 0 and 24 hours of incubation, the growth medium was removed, the cells were washed with PBS, and 500 µl of 0.25% sodium deoxycholate was added to lyse the infected macrophages. After 5 minutes the lysate was diluted in PBS, and the number of viable cells was determined after growth at 37° C. for 72 h on TSA plates. Duplicate samples were taken at all time points, and the assay was repeated two times.

#### Data Analyses

The student t test was performed in the analysis of colony forming units (cfu) data in the macrophage study (Lyman 25 Ott, 1988).

#### Results

Analysis of Nucleotide and Protein Sequence of *B. meliten-* sis tspA

The coding region of the tspA gene is 1329-bp long and is located between 219816 and 221144-bp on chromosome I of *B. melitensis* genome (accession number NC\_003317). The nucleotide sequence of the gene (SEQ ID NO: 7) is shown in FIG. 12A. The tspA gene is located immediately upstream of the gene encoding the invasion protein A of this chromosome. Analysis of the putative TspA protein sequence predicted that it contained a 24 amino acid long signal sequence (signal peptide probability: 0.999). The predicted cleavage site of the signal sequence is located between Gly24 and Ala25 residues (maximum cleavage site probability: 0.606). The theoretical mass of the TspA was 47-kDa. However, the predicted N-terminal processing of TspA would result in a mature protein having 419 amino acids and a molecular mass of 44.6-kDa. The amino acid sequence of the full length protein (SEQ ID NO: 8) is shown in FIG. 12B. The predicted subcellular localization of TspA was periplasmic space (Reliability Index: RI=1; Expected Accuracy=63%).

At the amino acid level, tspA gene shared 99% identity with the putative CtpA of B. suis. Additionally, it showed up to 77% identity with the putative CtpA proteins and periplasmic proteases of a number of bacterial species including Agrobacterium tumefaciens, Bartonella auintana. Bradyrhizobium japonicum, Magnetospirillum magnetotacticum, Mesorhizobium loti, Rhodobacter sphaeroides, Rhodopseudomonas palustris, and Sinorhizobium meliloti (data not shown). Part of the report showing the alignment of B. melitensis TspA with the closely related CtpA proteins and periplasmic proteases is shown in FIG. 13. Some motifs were conserved in all of the species analyzed, and the number of identical amino acid residues for the species included in FIG. 13 was 174 (data not shown). The amino acid residues that correspond with the amino acids critical for catalysis function of Synechocystis sp. strain PCC6803 CtpA protein (Inagaki et al., 2001) are indicated in FIG. 13. These residues were conserved among all the CtpA proteins and periplasmic proteases analyzed.

34 Discussion

A tspA mutant was generated by allelic exchange. A PCR amplification (see details in Experimental procedures) produced a predicted 1.4-kb size amplicon from the wild type *B. melitensis* strain 16M and an approximately 2.2-kb product from the tspA mutant strain 16MΔtspA, indicating that due to double-crossover event, a 471-bp region was deleted from the tspA gene, and the 1.3-kb Kan<sup>r</sup> was inserted at the deletion site of strain 16M genome. The nucleic acid sequence of the deletion mutant (SEQ ID NO: 9) and the corresponding amino acid sequence (SEQ ID NO: 10) are shown in FIGS. 14A and B, respectively. The deleted nucleotide sequence is shown in FIG. 14C (SEQ ID NO: 11). A number of PCR assays using the primers specific for a variety of *B. melitensis* genes confirmed that the tspA mutant was a *brucella* (data not shown).

Comparison of the Protein Profiles of B. melitensis Strains

The total protein profiles of the wild type, the tspA mutant, and the complemented tspA mutant strains were investigated by using denaturing gel electrophoresis and western immunoblotting. Two protein bands (approximately 28 and 65-kDa) were shown clearly upregulated in the tspA mutant strain. Those proteins were either absent or very faint in the wild type and the complemented strains.

#### Growth Rates of Recombinant B. melitensis Strains

The colonies of the tspA mutant appeared approximately half the size of the colonies of the wild type after three days of growth on TSA plates (data not shown). The wild type and the tspA mutant did not differ in terms of growth rates in LB broth (FIG. 15A). However, in contrast to the wild type, the tspA mutant exhibited a zero growth in salt-free LB broth (FIG. 15B). The colonies of the complemented tspA mutant appeared equal in size to those of the wild type (data not shown). The inability of the mutant strain to grow in salt-free broth was restored when this strain was complemented by introducing the *B. suis* ctpA gene (strain 16MΔtspA[ctpA<sup>+</sup>]). Additionally, the complemented strain grew at similar rates to the wild type strain in LB or salt-free LB broth (FIGS. 15A and 15B).

#### Cell Morphology

When examined by electron microscopy, the wild type strain grown in LB media displayed the native coccobacillus shape of *Brucella* (FIG. **16**A). In contrast, the tspA mutant strain grown in LB media acquired a spherical shape. <sup>45</sup> Additionally, some of the tspA mutant cells displayed a slightly to substantially increased cell diameter (FIG. **16**B).

Persistence of B. melitensis Strains in J774 Macrophages

The survival of *B. melitensis* strains in J774 mouse 50 macrophage cells was measured (Table 1). Relative to the survival of the wild type strain, that of the tspA mutant declined by  $0.2 \log_{10}$  cfu at 0 hours of incubation and  $1.2 \log_{10}$  cfu after 24 hours of incubation.

TABLE 1

Viability of B. melitensis strains in mouse macrophage J774 cells.

Recovery of *brucellae* from macrophages Mean ± SE log<sub>10</sub> cfu/well)<sup>a</sup>

Strain	0-hours of incubation	24-hours of incubation
16M	$5.72 \pm 0.07$	$4.84 \pm 0.09$
16M∆tspA	$5.52 \pm 0.04$	$3.65 \pm 0.07$

<sup>a</sup>P values for the difference between mean values were <0.07 for 0-hours of incubation and <0.005 for 24-hours of incubation

Based on its greater homology and substantial alignment with the bacterial carboxyl-terminal proteases and periplasmic proteases, we speculate that the putative TspA of B. melitensis is a carboxyl-terminal protease. The predicted signal sequence indicates that TspA is transported across the cytoplasmic membrane and thus suggests that this enzyme functions in the periplasmic space. Other reported bacterial carboxyl-terminal proteases like Tsp of E. coli (Hara et al., 1991) and CtpA of Borrelia burgdorferi (Ostberg et al., 2004) are also believed to function in the periplasmic space. The B. melitensis TspA amino acid sequence shared 99% homology with the B. suis CtpA sequence. The published protein sequence of B. suis CtpA (GenBank accession number NP 698817) does not include a signal sequence. This is because when the B. suis genomic sequence was annotated (Paulson et al., 2002), an 18 amino acid region had been mistakenly excluded from the N-terminal end of the actual CtpA amino acid sequence.

Thus far, the CtpA proteases have been well characterized only for the cyanobacterium Synechocystis sp. strain PCC 6803 (Anbudurai et al., 1994; Shestakov et al., 1994), the green alga S. obliquus (Trost et al., 1997), higher plants (Takahashi et al., 1988), and E. coli (Hara et al., 1989; 1991). A recent report by Ostberg et al., (2004) has partially characterized the substrates and the pleiotropic effects of CtpA protein of B. burgdorfei CtpA protein. In Synechocystis CtpA, among the conserved amino acids, Asp-253, Arg-255, Ser-313, Glu-316 and Lys-338 are critical for the in vivo activity (Inagaki et al., 2001). The B. burgdorferi CtpA protein and the E. coli TspA protein also carry the Asp, Arg, Ser, Glu and Lys residues corresponding to those residues of Synechocystis. Our sequence alignment work disclosed that the B. melitensis TspA sequence carries residues Asp238, Arg240, Ser300, Glu303, and Lys325 that correspond with those amino acid residues of CtpA sequences of Synechocystis, B. burgdorferi, and E. coli. Liao et al., (2000) reported that in CtpA protein of eukaryotic alga S. obliquus, the amino acid residues Ser372 and Lys397 form the catalytic center of this enzyme. These two amino acids are located 25 amino acids apart from each other in the CtpA proteins of Synechocystis species, S. obliquus, B. burgdorferi and E. coli. Consistent with these observations, in the B. melitensis TspA protein sequence, the Ser300 and Lys325 amino acids are located exactly 25 amino acids apart from each other. Accordingly, it could be postulated that these two amino acids make the Ser/Lys catalytic dyad of B. melitensis TspA.

By comparing the total protein expression profiles of the wild type and the tspA mutant B. melitensis strains, we found that expression of a number of proteins was altered by tspA inactivation. In fact the expression of at least two protein products (one of approximately 65-kDa and another of 28-kDa) was seen up-regulated due to mutation in the tspA gene. With the used immunoblotting procedure, it was impossible to detect if the expression of any other proteins was affected due to mutation. Further work using twodimensional gel electrophoresis is required to identify all the proteins that are affected by the tspA mutation. Ostberg et al., (2004) reported that the expression of the pore forming protein Oms28 was up-regulated when the ctpA gene encoding the CtpA protein was mutated in B. burgdorferi. These workers postulated the up-regulation of Oms28 protein as a result of a lack of C-terminal processing needed for regulating an inhibitor or an activator involved in modulation of Oms28 expression. Genetic complementation by introduction of a shuttle vector with the cloned ctpA gene showed restored protein expression in B. melitensis mutant to the

normal pattern (FIG. 3). Further experimentation will be required to determine whether this is a transcriptional or a posttranslational effect.

Western immunoblotting was not sensitive enough to identify the substrates of B. melitensis TspA protein. The protein D1 has been identified as the substrate of CtpA of cyanobacteria (Nixon et al., 1992), green algae (Liao et al., 2000) and higher plants (Takahashi et al., 1988). The PBP-3 and aberrant proteins are the substrates of E. coli Tsp (Hara et al., 1991, Keiler et al., 1996). The protein P13 and the hypothetical protein BB0323 have been identified as the potential targets of B. burgdorferi CtpA (Ostberg et al., 2004). The substrates of CtpA proteins of Brucella or any other bacteria have yet to be identified.

Just like the tsp mutant E. coli (Hara et al., 1991) and the ctpA mutant B. suis (Bandara et al., 2005, in press), the tspA mutant B. melitensis strain exhibited a zero growth in salt-free media suggesting that TspA function is involved either directly or indirectly in protection from osmotic stresses. Electron microscopy data revealed that similar to the ctpA mutant B. suis, the tspA mutant B. melitensis strain 20 possessed a spherical cell shape with some cells having increased cell diameter. Previous reports have shown that the E. coli strains deficient in penicillin-binding protein-2 (PBP-2) synthesis lacked the cell elongation pathway and grew as spherical cells because only septal synthesis was active in these strains (de Pedro et al., 2001). Similarly, a PBP-2 deficient strain of Erwinia amylovora displayed a large spherical phenotype, whereas its parent type counterpart displayed a rod-shaped phenotype (Milner et al., 1993). The diameter of the peptidoglycan stalk of Caulobacter crescentus increases when its PBP-2 homolog is inactivated (Seitz and Brun, 1998). These reports suggest that PBP-2 enzyme helps to regulate cellular diameter at the time of division. Based on these observations it can be hypothesized that the TspA protein is involved in processing of PBP-2 enzyme in B. melitensis, and disruption of tspA resulted in spherical shape cell with increased cell diameter.

In Bartonella bacilliformis genome, the gene encoding the CtpA is located immediately upstream of the gene encoding invasion associated locus A (ialA) (Mitchell and Minnick, 1997). Interestingly, in the B. melitensis genome, 40 Dalbey, R. E., and Von Heijne, G. (1992) Signal peptidases the tspA gene is located right upstream from the gene encoding the invasion protein A (locus tag BMEI0215, GeneID 1195927). Genetic complementation by introduction of a shuttle vector with the cloned tspA gene showed that the growth patterns were similar for the complemented tspA mutant and the wild type strain (FIG. 4B), and the total protein profiles were the same for the complemented strain and the wild type strain (FIG. 3). These complementation experiments confirmed that the phenotype observed after inactivation tspA gene was due to lack of expression of TspA alone and not due to a polar effect.

When introduced into macrophages, after 24 hours of incubation, the persistence of the tspA mutant strain declined significantly compared to that of the wild type strain indicating that TspA is important for survival, in particular against early killing by macrophages. Apparently, 55 due to the change of protein expression, and due to a possible influence on protein processing, the tspA mutant cells were unable to avoid phagosome-lysosome fusion, and thereby subjected to early killing.

In summary, the tspA gene of *B. melitensis* is homologous to the genes encoding carboxyl-terminal proteases and periplasmic proteases of other bacteria. Like the ctpA gene of B. bacilliformis, the tspA of B. melitensis is located immediately upstream of the gene encoding an invasion protein. Similar to the CtpA proteins of cyanobacteria, green algae, higher plants, E. coli, and B. burgdorferi, the TspA of B. melitensis contains all five amino acids critical for the CtpA catalytic activity, and a potential Ser/Lys catalytic dyad.

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Similar to the CtpA of B. burgdorferi, the TspA of B. melitensis influences the protein expression profile. Similar to the Tsp of E. coli and CtpA of B. suis, the TspA influences salt-sensitive growth and cell morphology of *B. melitensis*. In addition, similar to the CtpA of B. suis, the TspA of B. melitensis influences the intracellular persistence of the bacterium.

#### REFERENCES FOR EXAMPLE 3

- Acha, P., and Szyfres, B. (1980) Zoonoses and communicable diseases common to man and animals, p. 28-45. In Pan American Health Organization, Washington, D.C.
- Anbudurai P. R., Mor, T. S., Ohad, I., Shestakov, S, V, and Pakrasi, H. B. (1994) The ctpA gene encodes the C-terminal processing protease for the D1 protein of the photosystem II reaction center complex. Proc Natl Acad Sci USA 91:8082-8086.
- Banai, M., Adams, L. G., Frey, M., Pugh, R., and Ficht, T. A. (2002) The myth of Brucella L-forms and possible involvement of Brucella penicillin binding proteins in pathogenicity. Vet Microbiol 90:263-279.
- Bandara, A. B., Sriranganathan, N., Schurig, G. G. and Boyle, S. M. (2005) Putative Carboxyl-Terminal Protease Regulates Brucella suis Morphology in Culture and Persistence in Macrophages and Mice. J Bacteriol 187 (16) (In Press).
- Baumler A. J., Kusters J. G., Stojiljkovic I., and Heffron F. (1994) Salmonella typhimurium loci involved in survival within macrophages. Infect Immun 62:1623-1630.
- Bendtsen, J. D., Nielsen, H., von Heijne, G. and Brunak, S. (2004) Improved prediction of signal peptides: SignalP 3.0. J Mol Biol 340:783-795,
- 35 Corbel, M. J., and Brinley Morgan, W. J. (1984) Genus Brucella Meyer and Shaw 1920, 173 AL. p. 377-390. In N. R. Krieg and J. G. Holt (ed.), Bergey's Manual of Systematic Bacteriology, vol. 1. Williams and Wilkins, Baltimore.
- in prokaryotes and eukaryotes—a new protease family. Trends Biochem Sci 17:474-478.
- de Pedro, M. A., Donachie, W. D., Holtje, J. V., and Schwarz, H. (2001) Constitutive septal murein synthesis in Escherichia coli with impaired activity of the morphogenetic proteins RodA and penicillin-binding protein 2. J Bacteriol 183:4115-4126.
- Hara H., Nishimura, Y., Kato, J., Suzuki, H., Nagasawa, H., Suzuki, A., and Hirota, Y. (1989) Genetic analyses of processing involving C-terminal cleavage in penicillinbinding protein 3 of Escherichia coli. J Bacteriol 171: 5882-5889.
  - Hara, H., Yamamoto, Y., Higashitani, A., Suzuki, H., and Nishimura, Y. (1991) Cloning, mapping, and characterization of the Escherichia coli prc gene, which is involved in C-terminal processing of penicillin-binding protein 3. J Bacteriol 173:4799-4813.
- Inagaki, N., Maitra, R., Satoh, K., Pakrasi, H. B. (2001) Amino acid residues that are critical for in vivo catalytic activity of CtpA, the carboxyl-terminal processing protease for the D1 protein of photosystem II. J Biol Chem 276:30099-30105.
- Inagaki, N., Yamamoto, Y., Mori, H., and Satoh, K. (1996) Carboxyl-terminal processing protease for the D1 precursor protein: cloning and sequencing of the spinach cDNA. Plant Mol Biol 30:39-50.

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- Keiler, K. C., Waller, P. R. H., and Sauer, R. T. (1996) Role of a peptide tagging system in degradation of proteins synthesized from damaged messenger RNA. *Science* 271: 990-993.
- Kovach, M. E., Phillips, R. W., Elzer, P. H., Roop II R. M., 5 and Peterson, K. M. (1994) pBBR1 MCS: a broad-host range cloning vector. *Bio Techniques* 16:800-802.
- Laemmli, U. K. (1970) Cleavage of structural proteins during the assembly of the head of bacteriophage T4. *Nature* 227:680-685.
- Liao, D.-I., Qian, J., Chisholm, D. A., Jordan, D. B., and Diner, B. A. (2000) *Nat Struct Biol* 7:749-753.
- Lyman Ott, R. (1988) An Introduction to Statistical Methods and Data Analysis. Fourth Edition. Duxbury Press. Belmont, Calif.
- McQuiston, J. R., Schurig, G. G., Sriranganathan, N., and Boyle S. M. (1995) Transformation of *Brucella* species with suicide and broad host-range plasmids. *Methods Mol Bio* 47:143-148. Milner, J. S., Dymock, D., Cooper, R. M., and Roberts, I. S. (1993) Penicillin-binding proteins from *Erwinia amylovora*: mutants lacking PBP2 are avirulent. *J Bacteriol* 175:6082-6088. Mitchell S. J., and Minnick, M. F. (1997) A carboxy-terminal processing protease gene is located immediately upstream of the invasion-associated locus from *Bartonella bacilliformis*. 25 *Microbiology* 143:1221-33.
- Nixon P. J., Trost J. T., and Diner B. A. (1992) Role of the carboxy terminus of polypeptide D1 in the assembly of a functional wateroxidizing manganese cluster in photosystem II of the cyanobacterium *Synechocystis* sp. PCC 30 6803: assembly requires a free carboxyl group at C-terminal position 344. *Biochemistry* 31:1085910871.
- Oelmüller, R., Herrmann, R. G. and Pakrasi, H. B. (1996) Molecular studies of CtpA, the carboxyl-terminal processing protease for the D1 protein of the photosystem II 35 reaction center in higher plants. *J Biol Chem* 271:21848-21856.
- Ostberg, Y., Carroll, J. A., Pinne, M., Krum, J. G., Rosa, P., and Bergstrom, S. (2004) Pleitropic effects of inactivating a carboxyl-terminal protease CtpA, in *Borrelia burgdor-* 40 *feri. J Bacteriol* 186:2074-2084.
- Paetzel, M. and Dalbey, R. E. (1997) Catalytic hydroxyl/amine dyads within serine proteases. *Trends Biochem Sci* 22:28-31. Paulson, I. T., Sheshadri R., Nelson K. E. et al., (2002) The *Brucella suis* genome reveals fundamental 45 similarities between animal and plant pathigens and symbionts. *Proc Natl Acad Sci USA* 99:13148-13153.
- Popham, D. L., and Young, K. D. (2003) Role of penicillinbinding proteins in bacterial cell morphogenesis. *Current Opinion in Microbiology* 6:594-599.
- Roop, R. M. 2 nd, Price, M. L., Dunn, B. E., Boyle, S. M, Sriranganathan, N., and Schurig, G. G. (1992) Molecular

- cloning and nucleotide sequence analysis of the gene encoding the immunoreactive *Brucella abortus* Hsp60 protein, BA60 K. *Microb Pathog* 12:47-62.
- Schurig, G. G., Roop, R. M., Bagchi, T., Boyle, S. M., Buhrman, D., and Sriranganathan, N. (1991) Biological properties of RB51: a stable rough strain of *Brucella* abortus. Vet Microbiol 28:171-188.
- Seitz, L. C., and Brun, Y. V. (1998) Genetic analysis of mecillinam-resistant mutants of *Caulobacter crescentus* deficient in stalk biosynthesis. *J Bacteriol* 180:5235-5239
- Shestakov, S. V., Anbudurai, P. R., Stanbekovas, G. E., Gadzhiev, A., Lind L. K., and Pakrasi, H. B. (1994) Molecular cloning and characterization of the ctpA gene encoding a carboxyl-terminal processing protease. Analysis of a spontaneous photosystem II-deficient mutant strain of the cyanobacterium *Synechocystis* sp. PCC 6803. *J Biol Chem* 269:19354-19359.
- Silber, K. R., Keiler, K. C., and Sauer, R. T. (1992) Tsp: a tail-specific protease that selectively degrades proteins with nonpolar C termini. *Proc Natl Acad Sci U.S.A* 89:295-299.
- Takahashi, M., Shiraishi, T., and Asada, K. (1988) COOH-terminal residues of D1 and the 44 kDa CPa-2 at spinach photosystem II core complex. *FEBS Lett* 240:6-8.
- Trost, J. T., Chisholm, D. A., Jordan, D. B., and Diner, B. A. (1997) The D1 C-terminal processing protease of photosystem II from *Scenedesmus obliquus*. Protein purification and gene characterization in wild type and processing mutants. *J Biol Chem* 272:20348-20356.
- Vemulapalli, R., Duncan, A. J., Boyle, S. M., Sriranganathan, N., Toth, T. E., and Schurig, G. G. (1998) Cloning and sequencing of yajC and secD homologs of *Brucella abortus* and demonstration of immune responses to YajC in mice vaccinated with *B. abortus* RB51. *Infect Immun* 66:5684-5691.
- Yamamoto, Y., Inagaki, N., and Satoh, K. (2001) Overexpression and characterization of carboxyl-terminal processing protease for precursor D1 protein: regulation of enzyme-substrate interaction by molecular environments. *J Biol Chem* 276:7518-7525
- Young, E. J. (1983) Human brucellosis. Rev Infect Dis 5:821-842.
- While the invention has been described in terms of its preferred embodiments, those skilled in the art will recognize that the invention can be practiced with modification within the spirit and scope of the appended claims. Accordingly, the present invention should not be limited to the embodiments as described above, but should further include all modifications and equivalents thereof within the spirit and scope of the description provided herein.

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Leu Thr Ser	Leu As 20	o Pro	His	Ser	Ser 25	Tyr	Met	Asn	Ser	Thr 30	Asp	Ala
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Glu Val Thr 50  Asp Thr Pro 65  Lys Ile Asp Asp Lys Met  Arg Lys Gly 115  Ile Ala Val 130  Leu Arg Val 145  Gly Ile Glu	Met Gl' Ala Al Gly Gl' 85 Arg Gl' 100 Ala Gl' Arg Al Ile Se Lys Il 16 Asp Le 180	a Arg 70 h Asp Ala Lys a Val r Phe 150 c Gln i Arg	Asp 55 Ala Val Val Pro Lys 135 Thr Ala	40 Leu Gly Arg Gly Ile 120 Val Glu Asn	Val Gly Thr 105 Glu Arg Lys Val Pro 185	Leu Leu 90 Pro Leu Val Thr Pro 170 Gly	Val Ala 75 Lys Ile Thr Glu Phe 155 Ala Gly	Thr 60 Gly Leu Lys Ile Gly 140 Glu Asp Leu	Asp Glu Leu Val 125 Asp Lys Lys	Pro Phe Glu Thr 110 Arg Val Leu Leu Asp 190	Ile Ile Ala 95 Ile Asp Gly Lys 175 Gln	Asp Ser 80 Val Leu Val Tyr Lys 160 Gly Ala

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Tyr	Tyr 290	Thr	Pro	Ser	Gly	Lys 295	Ser								

### We claim:

1. An attenuated, recombinant *Brucella melitensis* strain deficient in the carboxyl-terminal proteolytic processing activity of a tail-specific protease wherein said deficiency is caused by a nucleotide sequence deletion within a tspA gene that, absent said nucleotide sequence deletion, encodes the amino acid sequence of the TspA protease set forth in SEQ ID NO:8.

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- 2. The attenuated, recombinant *Brucella melitensis* strain of claim 1, wherein said attenuated recombinant *Brucella melitensis* strain is  $16\text{M}\Delta \text{tspA}$ .
- 3. A method for eliciting an immune response to *Brucella melitensis* in a mammal, or treating Brucellosis in a mammal comprising administering to said mammal an attenuated, recombinant, *Brucella melitensis* strain deficient in the carboxyl-terminal proteolytic processing activity of a tail-specific protease wherein said deficiency is caused by a nucleotide sequence deletion within a tspA gene that, absent said nucleotide sequence deletion, encodes the amino acid sequence of the TspA protease set forth in SEQ ID NO:8,

- wherein said attenuated, recombinant, *Brucella melitensis* strain is administered in a quantity sufficient to elicit an immune response, thereby eliciting an immune response.
- **4**. The method of claim **3**, wherein said attenuated, recombinant *Brucella melitensis* strain is 16MΔtspA.
- 5. The method of claim 3, wherein said mammal is selected from the group consisting of humans, sheep, goats, dogs, swine, cattle, and reindeer.
- **6**. A composition for eliciting an immune response to *Brucella melitensis* in a mammal, comprising
  - (i) an attenuated, recombinant, *Brucella melitensis* strain with a deficiency in the carboxypeptidase activity of a tail-specific protease, wherein said deficiency is caused by a nucleotide sequence deletion in a tspA gene that, absent said nucleotide sequence deletion, encodes the TspA carboxypeptidase having the amino acid sequence set forth in SEQ ID NO:8; and
- (ii) a physiologically suitable carrier.
- 7. The composition of claim 6, wherein said attenuated, recombinant *Brucella melitensis* strain is  $16\text{M}\Delta \text{tspA}$ .

\* \* \* \* \*

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